SUBSTITUTED TRICYCLIC GAMMA-CARBOLINES AS SEROTONIN RECEPTOR AGONISTS AND ANTAGONISTS

RELATED APPLICATIONS

This application claims priority benefit under Title 35 § 119(e) of United States provisional Application No. 60/434,760, filed December 19, 2002, the contents of which are herein incorporated by reference.

FIELD OF THE INVENTION

The present invention is directed to novel compounds represented by structural Formula (I):

$$\begin{array}{c|c}
R^9 & & \\
R^8 & & \\
\hline
R^7 & & \\
R^6 & & \\
\hline
R^5 & \\
\hline
(I)
\end{array}$$

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or a pharmaceutically acceptable salt thereof, wherein R¹, R^{4a}, R⁵, R⁶, R⁷, R⁸, R⁹, and m, are defined herein. The invention is also concerned with pharmaceutical formulations comprising these novel compounds as active ingredients and the use of the novel compounds and their formulations in the treatment of certain central nervous system disorders. The compounds of this invention are serotonin receptor modulators, in particular 5HT_{2C} receptor agonists and antagonists, and are useful in the control or prevention of central nervous system disorders including obesity, anorexia, bulemia, depression, anxiety, psychosis, schizophrenia, migraine, addictive behavior, obsessive-compulsive disorder, and sexual disorders.

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BACKGROUND OF THE INVENTION

The neurotransmitter/hormone serotonin (5-hydroxytryptamine, 5-HT) regulates many physiological processes via a group of at least 14 distinct receptors that are organized into 7 subfamilies (Hoyer, D., et al., Pharmacol. Rev., 46, 1994). The 5-HT₂ subfamily is composed of the 5-HT_{2A}, 5-HT_{2B}, and 5-HT_{2C} receptors as determined by gene homology and pharmacological properties. There exists a substantial correlation for the relationship between 5-HT₂ receptor modulation and a variety of diseases and therapies. Prior to the early 1990's the 5-HT_{2C} and 5-HT_{2A} receptors were referred to as 5-HT_{1C} and 5-HT₂, respectively.

The direct or indirect agonism or antagonism of 5-HT₂ receptors, either selectively or non-selectively, has been associated with the treatment of various central nervous system (CNS) disorders including obesity, depression, schizophrenia and bi-polar disorders. In the recent past the contribution of serotonergic activity to the mode of action of anti-obesity drugs has been well documented. Compounds that increase the overall basal tone of serotonin in the CNS have been successfully developed as anorectic drugs. The serotonin releasing agents, such as fenfluramine, function by increasing the amount of serotonin present in the nerve synapse. These breakthrough treatments, however, are not without side effects. Due to the mechanism of action of serotonin releasing agents, they effect the activity of a number of serotonin receptor subtypes in a wide variety of organs including those not associated with the desired mechanism of action. This non-specific modulation of the serotonin family of receptors most likely plays a significant role in the side effect profile. In addition, these compounds or their metabolites often have a high affinity for a number of the serotonin receptors as well as a multitude of other monoamine neurotransmitters and nuisance receptors. Removing some of the receptor cross reactivity would allow for the examination and possible development of potent therapeutic ligands with an improved side effect profile.

The 5-HT_{2C} receptor is a G-protein coupled receptor. It is almost exclusively expressed in the central nervous system including the hypothalamus, hippocampus, amygdala, nucleus of the solitary tract, spinal cord, cortex, olfactory bulb, ventral tegmental area (VTA), nucleus accumbens and choroid plexus (Hoffman, B. and Mezey, E., FEBS Lett., 247, 1989). There is ample evidence to support the role of

selective 5-HT_{2C} receptor ligands in a number of disease therapies. 5-HT_{2C} knockout mice develop a late stage obesity syndrome that is not reversed by fenfluramine or other direct acting 5-HT_{2C} agonists such as mCPP (Nonogaki, K., et al., Nature Med., 4, 1998; Vickers, S., et. al., Psychopharmacology, 143, 1999). Administration of 5 selective 5-HT_{2C} agonists to rats causes a reduction in food intake and corresponding reduction in body weight (Vickers, S., et al., Br. J. Pharmacol., 130, 2000) and these responses can be blocked by administration of selective 5-HT_{2C} antagonists (Vicker, S., et al., Neuropharmacol., 41, 2001). 5-HT_{2C} receptor modulation in the hypothalamus can also influence thermoregulation (Mazzola-Pomietto, P, et al., 10 Psychopharmacology, 123, 1996), sleep (Sharpley, A., et al., Neuropharmacology, 33, 1994), sexual behavior and neuroendocrine function (Rittenhouse, P. et al., J. Pharmacol. Exp. Ther., 271, 1994). Activation of 5-HT_{2C} receptors in the VTA modulates the activity of dopaminergic neurons that are involved in aspects of depression (Di Matteo, V. et al., Trends Pharmacol. Sci., 22, 2001) and 5-HT_{2C} 15 receptor agonists such as WAY 161503, RO 60-0175 and RO 60-0332 are active in rodent models of depression (Cryan, J. and Lucki, I., J. Pharmacol. Exp. Ther., 295, 2000). 5-HT_{2C} agonists have been reported to reduce the rewarding effects of nicotine administration in rats (Grottick, A., et al., Psychopharmacology, 157, 2001) and influences rodent responses to cocaine administration (Grottick, A., et al., J. 20 Pharmacol. Exp. Ther., 295, 2000). Modulation of 5-HT_{2C} receptors in the spinal cord can influence pain perception (Chojnacka-Wojcik, E., et al., Pol. J. Pharmacol., 46, 1994). There is also data indicating that the 5-HT_{2C} receptor agonists mCPP and RO 60-0175 mediate penile erections in rats (Millan, M., et al., Eur J. Pharmacol. 325, 1997).

Compounds reported to bind to and activate 5-HT_{2C} receptors are disclosed in the following documents. U.S. Patent Numbers 3,914,421; 4,013,652; 4,115,577; 4,183,936; and 4,238,607 disclose pyridopyrrolobenz-heterocycles of formula:

where X is O, S, S(=O), or SO₂; n is 0 or 1; R^1 is various carbon substituents, and Z is a monosubstituent of H, methyl, or chloro.

U.S. Patent Number 4,219,550 discloses pyridopyrrolo-benzheterocycles of formula:

where X is O or S; R^1 is C_{1-4} alkyl or cyclopropyl; R^2 is H, CH₃, OCH₃, Cl, Br, F, or CF₃; and (A) is -CH₂-, -CH(CH₃)-, or -CH₂CH₂-.

European Patent Application EP 473,550 A1 discloses indolonaphthyridines of formula:

$$R^3$$
 R^4
 R^5

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wherein X and Y are H or a simple ring, R^1 , is H, alkyl, alkylcarbonylalkyl, arylcarbonylalkyl, aralkyl, or a mono or disubstituted carbamoylalkyl; and R^3 , R^4 , and R^5 are H, halogen, alkyl, alkoxy, alkylthio or trifluoromethyl.

PCT International Patent Application WO 00/35922 discloses tetrahydro-1H-pyrazino(1,2-A-quinoxalin-5(6H)one derivatives of formula:

$$R^2$$
 R^3
 R^4
 R^4
 R^7

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as being $5HT_{2C}$ agonists; wherein X is CR^5R^6 or carbonyl; R is H or alkyl; R' is H, alkyl, acyl, or aroyl; and R^1 , R^2 , R^3 , and R^4 are independently, H, alkyl, alkoxy, halogen, trifluoroalkyl, cyano, alkylsulfonamide, alkyl amide, amino, alkylamino, dialkylamino, trifluoroalkoxy, acyl, or aryl.

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U.S. Patent Number 6,552,017, 6,548,493 PCT International Patent Application WO 00/77001, WO 00/77002, and WO 00/77010 discloses substituted heterocycle fused gamma-carbolines of formula:

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wherein X is CHR, C(=O), O, S, S(=O), SO₂, NR, C(=O)NR, or NRC(=O); n is 0, 1 or 2; m is 0, 1, 2 or 3; k is 1 or 2: R¹ is H, alkyl, alkenyl, alkynyl, cycloalkyl, acyl, or aryl; R⁵ is H or alkyl, R^{6a} and R^{6b} are independently H, OH, alkyl, alkenyl, alkynyl, cycloalkyl, alkoxy, trifluoroalkyl, alkylamino, trifluoroalkoxy, acyl, or aryl; and R⁷, R⁸, and R⁹ are independently, H, alkyl, alkenyl, alkynyl, cycloalkyl, alkoxy, halogen, trifluoroalkyl, cyano, alkylsulfonamide, alkyl amide, amino, alkylamino, dialkylamino, trifluoroalkoxy, acyl, aryl, or heterocyclic ring.

Patent Application WO 02/59129 discloses substituted pyridoindoles of formula:

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wherein X is O, S, S(=O), SO₂, or NR; n is 1 or 2; k is 1 or 2: R¹ is H, alkyl, alkenyl, alkynyl, cycloalkyl, acyl, or aryl; R⁵ and R⁶ are independently H, or alkyl: and R⁷, R⁸, and R⁹ are independently, H, alkyl, alkenyl, alkynyl, cycloalkyl, alkoxy, halogen, trifluoroalkyl, cyano, alkylsulfonamide, alkyl amide, amino, alkylamino, dialkylamino, trifluoroalkoxy, acyl, aryl, or heterocyclic ring.

PTC International Patent Application WO 03/14118 discloses 1H-pyrido[4,3-b]indoles of formula:

$$R^2$$
 R^3
 R^4
 R^5

wherein R^1 , R^2 , R^3 and R^4 are independently, H, halo, CF3, OCF3, CN, NO2, alkyl, alkenyl, alkynyl, cycloalkyl, cycloalkenyl, alkoxy, thioalkyl, C(=O)Ar, aryl, or alkyleneAr, provided that at least one of R^1 , R^2 , R^3 or R^4 is aryl; and R^5 , R^6 and R^7 are independently H, or various carbon substituents.

None of the above references suggest or disclose the compounds of the present invention. There remains a need to discover new compounds useful as serotonin receptor modulators, i.e. selective agonists and antagonists, which are useful in the control or prevention of central nervous system disorders. As such, the present invention discloses novel compounds which are useful as serotonin agonists and antagonists, and provide good in vitro potency.

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SUMMARY OF THE INVENTION

One object of the present invention is to provide novel compounds which are useful as agonists of the 5-HT2C receptor, or pharmaceutically acceptable salts or prodrugs thereof.

It is another object of the present invention to provide pharmaceutical compositions comprising a pharmaceutically acceptable carrier and a therapeutically effective amount of at least one of the compounds of the present invention or a pharmaceutically acceptable salt or prodrug form thereof.

It is another object of the present invention to provide a method for treating central nervous system disorders including obesity, anorexia, bulemia, depression, anxiety, psychosis, schizophrenia, migraine, addictive behavior, obsessive-compulsive disorder, and sexual disorders comprising administering to a host in need of such treatment a therapeutically effective amount of at least one of the compounds of the present invention or a pharmaceutically acceptable salt or prodrug form thereof. More specifically, the present invention provides a method for treating obesity.

These and other objects, which will become apparent during the following detailed description, have been achieved by the inventors' discovery that compounds of Formula (I):

$$\begin{array}{c|c}
R^9 & & \\
R^8 & & \\
\hline
R^7 & & \\
R^6 & & \\
\hline
R^5 & & \\
\hline
(I)
\end{array}$$

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or pharmaceutically acceptable salt or prodrug forms thereof, wherein R¹, R^{4a}, R⁵, R⁶, R⁷, R⁸, R⁹ and m are defined below, are effective agonists of the 5-HT2C receptor.

DETAILED DESCRIPTION OF THE EMBODIMENTS

Thus, in a first embodiment, the present invention provides a novel compound of Formula (I):

$$\begin{array}{c|c}
R^{9} & & \\
R^{8} & & \\
\hline
R^{7} & & \\
R^{6} & & \\
\hline
R^{5} & & \\
\end{array}$$
(I)

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or a stereoisomer or a pharmaceutically acceptable salt form thereof, wherein:

10 R¹ is selected from

H, $C(=O)R^{2a}$, $C(=O)OR^{2a}$, $S(=O)R^{2a}$, $S(=O)_2R^{2a}$,

C3-7 cycloalkyl,

C₁₋₄ alkyl substituted with 0-3 R²,

C₂₋₄ alkenyl substituted with 0-2 R²,

15 C₂₋₄ alkynyl substituted with 0-2 R²,

aryl substituted with 0-5 R⁴²;

C₃₋₁₀ carbocyclic residue substituted with 0-3 R⁴¹, and

5-6 membered heterocyclic ring system containing from 1-4 heteroatoms selected from the group consisting of N, O, and S substituted with 0-3

 R^{41} ;

R², at each occurrence, is independently selected from

halo, C₁₋₃ haloalkyl, C₁₋₄ alkoxy, C₁₋₄ alkyl,

C2-4 alkenyl, C2-4 alkynyl, C3-6 cycloalkyl,

25 aryl substituted with 0-5 R⁴²;

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C<sub>3-10</sub> carbocyclic residue substituted with 0-3 R<sup>41</sup>, and
                   5-6 membered heterocyclic ring system containing from 1-4 heteroatoms
                            selected from the group consisting of N, O, and S substituted with 0-3
                            R41:
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        R^{2a} is H, C<sub>1-4</sub> alkyl, (aryl)C<sub>1-4</sub> alkyl-, or
                   (C<sub>3-6</sub> cycloalkyl)C<sub>1-4</sub> alkyl-;
        R<sup>4a</sup> is H or C<sub>1-4</sub> alkyl;
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        R<sup>5</sup> is H, C<sub>1-4</sub> alkyl substituted with 0-2 R<sup>20</sup>,
                  -C(=O)(C_{1-4} \text{ alkyl}), -C(=O)O(C_{1-4} \text{ alkyl}), or C_{1-4} \text{ haloalkyl};
        R<sup>6</sup> is selected from
                  halo, -CF3, -OCF3, -CN, -NO2, -OCH3, -SCH3, -CF2CF3, -O-R<sup>11</sup>,
15
                  -OCF2CF3, -OCF2H, -OCF2CH3,
                  -S-R^{11}, -S(=O)-R^{11}, -S(=O)2-R^{11}, -S(=O)-NR^{10}-R^{11},
                  -S(=O)2-NR<sup>10</sup>-R<sup>11</sup>, -NR<sup>10</sup>-R<sup>11</sup>, -CH<sub>2</sub>O-R<sup>11</sup>, -CH<sub>2</sub>S-R<sup>11</sup>,
                  -CH_2S(=O)-R^{11}, -CH_2S(=O) 2-R<sup>11</sup>, -CH_2NR^{10}-R^{11}, -C(=O)NR^{10}-R^{11}
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                  C<sub>1-4</sub> haloalkyl, (C<sub>1-4</sub> haloalkyl)oxy;
                  C<sub>1-4</sub> alkyl substituted with 0-2 R<sup>20</sup>,
                  C<sub>2-4</sub> alkenyl substituted with 0-2 R<sup>20</sup>,
                  C<sub>2-4</sub> alkynyl substituted with 0-1 R<sup>20</sup>, and
                  C<sub>3-6</sub> carbocyclic residue substituted with 0-3 R<sup>21</sup>,
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        R<sup>7</sup> and R<sup>9</sup> are independently selected from
                  H, F, Cl, Br, -CF3, -OCF3, -OH, -CN, -NO2, -CF2CF3, C1-4 alkyl,
                  C2-4 alkenyl, C2-4 alkynyl, C1-4 haloalkyl, C1-4 alkoxy, and
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(C₁₋₄ haloalkyl)oxy;

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R<sup>8</sup> is selected from
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halo, -CF3, -OCF3, -OH, -CN, -NO2, -OCH3, -SCH3, -CF2CF3,

5 $-OR^{12}$, $-SR^{12}$, $-NR^{12}R^{13}$, -C(O)H, $-C(O)R^{12}$, $-C(O)NR^{12}R^{13}$,

 $-NR^{14}C(O)R^{12}$, $-C(O)OR^{12}$, $-OC(O)R^{12}$, $-OC(O)OR^{12}$,

 $-S(O)R^{12}$, $-S(O)_2R^{12}$, $-S(O)NR^{12}R^{13}$, $-S(O)_2NR^{12}R^{13}$,

 $-NR^{14}S(O)R^{12}$, $-NR^{14}S(O)_2R^{12}$, $-NR^{12}C(O)R^{15}$, $-NR^{12}C(O)OR^{15}$,

-NR¹²S(O)₂R¹⁵, -NR¹²C(O)NHR¹⁵;

10 C₁₋₆ alkyl substituted with 0-2 R^{8a},

C₂₋₆ alkenyl substituted with 0-2 R^{8a},

C₂₋₆ alkynyl substituted with 0-2 R^{8a},

C₃₋₆ cycloalkyl substituted with 0-2 R^{8a},

C₃₋₁₀ carbocyclic residue substituted with 0-3 R³³;

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 R^{8a} , at each occurrence, is independently selected from

halo, -CF3, -OCF3, -OH, -CN, -NO2, -CF2CF3,

methyl, ethyl, n-propyl, i-propyl, n-butyl, i-butyl, s-butyl, t-butyl,

 $-OR^{12}$, $-SR^{12}$, $-NR^{12}R^{13}$, -C(O)H, $-C(O)R^{12}$, $-C(O)NR^{12}R^{13}$,

20 $-NR^{14}C(O)R^{12}$, $-C(O)OR^{12}$, $-OC(O)R^{12}$, $-OC(O)OR^{12}$,

 $-S(O)R^{12}$, $-S(O)_2R^{12}$, $-S(O)NR^{12}R^{13}$, $-S(O)_2NR^{12}R^{13}$,

 $-NR^{14}S(O)R^{12}$, $-NR^{14}S(O)_2R^{12}$, $-NR^{12}C(O)R^{15}$, $-NR^{12}C(O)OR^{15}$,

-NR¹²S(O)₂R¹⁵, -NR¹²C(O)NHR¹⁵;

phenyl substituted with 0-5 R³³;

25 C₃₋₁₀ carbocyclic residue substituted with 0-3 R³³, and

5-10 membered heterocyclic ring system containing from 1-4 heteroatoms selected from the group consisting of N, O, and S substituted with 0-3 R^{33} ;

R¹⁰ is H or C₁₋₄ alkyl;

R¹¹ is selected from

5 C_{1-6} alkyl substituted with 0-2 R^{20} ,

C₂₋₆ alkenyl substituted with 0-2 R²⁰,

C₂₋₆ alkynyl substituted with 0-1 R²⁰,

C₃₋₁₀ carbocyclic residue substituted with 0-3 R²¹,

aryl substituted with 0-5 R²³, and

5-10 membered heterocyclic ring system containing from 1-4 heteroatoms selected from the group consisting of N, O, and S substituted with 0-3 R²¹;

alternatively, R^{10} and R^{11} join to form a 5- or 6-membered ring optionally substituted with -O- or -N(R^{14})-;

alternatively, R¹⁰ and R¹¹ when attached to N may be combined to form a 9- or 10membered bicyclic heterocyclic ring system containing from 1-3 heteroatoms
selected from the group consisting of N, O, and S, wherein said bicyclic
heterocyclic ring system is unsaturated or partially saturated, wherein said
bicyclic heterocyclic ring system is substituted with 0-3 R¹⁶;

R¹² is selected from H,

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C₁₋₆ alkyl substituted with 0-2 R^{12a},

C₂₋₆ alkenyl substituted with 0-2 R^{12a},

C₂₋₆ alkynyl substituted with 0-2 R^{12a},

C₃₋₆ cycloalkyl substituted with 0-3 R³³,

aryl substituted with 0-5 R³³;

 C_{3-10} carbocyclic residue substituted with 0-3 R^{33} , and

- 5-10 membered heterocyclic ring system containing from 1-4 heteroatoms selected from the group consisting of N, O, and S substituted with 0-3 R³³:
- 5 R^{12a}, at each occurrence, is independently selected from

$$-SO_2NR^{46}R^{47}$$
, $-CONR^{46}R^{47}$, $-OR^{45}$, $=O$,

C₁₋₄ alkyl, C₂₋₆ alkenyl, C₂₋₆ alkynyl,

phenyl substituted with 0-5 R³³;

C₃₋₁₀ carbocyclic residue substituted with 0-3 R³³, and

5-10 membered heterocyclic ring system containing from 1-4 heteroatoms selected from the group consisting of N, O, and S substituted with 0-3 R^{33} ;

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- R¹³, at each occurrence, is independently selected from H, C₁₄ alkyl, C₂₄ alkenyl, and C₂₄ alkynyl;
- alternatively, R^{12} and R^{13} join to form a 5- or 6-membered ring optionally substituted with -O- or -N(R^{14})-;
- alternatively, R¹² and R¹³ when attached to N may be combined to form a 9- or 10membered bicyclic heterocyclic ring system containing from 1-3 heteroatoms
 selected from the group consisting of N, O, and S, wherein said bicyclic
 heterocyclic ring system is unsaturated or partially saturated, wherein said
 bicyclic heterocyclic ring system is substituted with 0-3 R¹⁶;
 - R¹⁴, at each occurrence, is independently selected from H and C₁₋₄ alkyl;
- 30 R¹⁵, at each occurrence, is independently selected from

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H, C₁₋₄ alkyl, C₂₋₄ alkenyl, and C₂₋₄ alkynyl;

 R^{16} , at each occurrence, is independently selected from

H, OH, halo, CN, NO₂, CF₃, SO₂R⁴⁵, NR⁴⁶R⁴⁷, -C(=O)H,

C₁₋₄ alkyl, C₂₋₄ alkenyl, C₂₋₄ alkynyl, C₁₋₄ haloalkyl,

C₁₋₃ haloalkyl-oxy-, and C₁₋₃ alkyloxy-;

R²⁰ is selected from

H, halo, -OH, -CF3, -CN, -NO2, -CO2H, -SO2R⁴⁵, -SOR⁴⁵, -SR⁴⁵,

10 -NR⁴⁶SO₂R⁴⁵, -NR⁴⁶COR⁴⁵, -NR⁴⁶R⁴⁷,

C₁₋₄ alkyl, C₂₋₆ alkenyl, C₂₋₆ alkynyl, C₁₋₄ alkoxy,

C₁₋₄ haloalkyl;

C₃₋₁₀ carbocyclic residue substituted with 0-3 R²¹;

aryl substituted with 0-5 R²³; and

- 5-10 membered heterocyclic ring system containing from 1-4 heteroatoms selected from the group consisting of N, O, and S substituted with 0-3 R²¹;
- R²¹, at each occurrence, is independently selected from

 H, OH, halo, CF₃, SO₂R⁴⁵, NR⁴⁶R⁴⁷, CN, NO₂, =O, C₁₋₄ alkyl,

 C₁₋₄ alkoxy, and (C₁₋₄ haloalkyl)oxy;
 - R²³, at each occurrence, is independently selected from
 H, OH, halo, CF₃, SO₂R⁴⁵, NR⁴⁶R⁴⁷, CN, NO₂, C₁₋₄ alkyl,
 C₁₋₄ alkoxy, and (C₁₋₄ haloalkyl)oxy;
 - R^{33} , at each occurrence, is independently selected from H, OH, halo, -CN, -NO₂, -CF₃, -OCF₃, -SO₂ R^{35} , -S(=O) R^{35} , -SR³⁵, -NR³⁶ R^{37} , -NHC(=O) R^{35} , -C(=O)NR³⁶ R^{37} .

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-C(=O)H, -C(=O)R^{35}, -C(=O)OR^{35}, -OC(=O)R^{35}, -OR^{35}.
                     C<sub>1-6</sub> alkyl, C<sub>2-6</sub> alkenyl, C<sub>2-6</sub> alkynyl, C<sub>1-4</sub> haloalkyl,
                     C<sub>1-4</sub> alkoxy, (C<sub>1-4</sub> haloalkyl)oxy,
                     C<sub>3-6</sub> cycloalkyl, phenyl, aryl substituted with 0-2 R<sup>34</sup>,
                     C<sub>1-6</sub> alkyl substituted with R<sup>34</sup>, and
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                     C<sub>2-6</sub> alkenyl substituted with R<sup>34</sup>;
         R<sup>34</sup>, at each occurrence, is independently selected from
                     OH, C_{1-4} alkoxy, -SO_2R^{35}, -NR^{36}R^{37}, NR^{36}R^{37}C(=0)-, and
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                     (C<sub>1-4</sub> alkyl)CO<sub>2</sub>-;
         R<sup>35</sup>, at each occurrence, is independently selected from
                     C<sub>1-4</sub> alkyl, C<sub>1-4</sub> haloalkyl, C<sub>3-6</sub> cycloalkyl,
                    (C<sub>3-6</sub> cycloalkyl)methyl-, and (C<sub>3-6</sub> cycloalkyl)ethyl-;
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         R<sup>36</sup>, at each occurrence, is independently selected from H and C<sub>1-4</sub> alkyl;
         R<sup>37</sup>, at each occurrence, is independently selected from H, C<sub>1-4</sub> alkyl,
                    -C(=O)NH(C_{1-4} \text{ alkyl}), -SO_2(C_{1-4} \text{ alkyl}),
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                    -C(=O)O(C_{1-4} \text{ alkyl}), -C(=O)(C_{1-4} \text{ alkyl}), \text{ and } -C(=O)H;
         R<sup>41</sup>, at each occurrence, is independently selected from
                    H, CF<sub>3</sub>, halo, OH, CO<sub>2</sub>H, SO<sub>2</sub>R<sup>45</sup>, NR<sup>46</sup>R<sup>47</sup>, NO<sub>2</sub>, CN, =O,
                    C<sub>1-4</sub> alkyl, C<sub>2-8</sub> alkenyl, C<sub>2-8</sub> alkynyl, C<sub>1-4</sub> alkoxy, and C<sub>1-4</sub> haloalkyl;
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         R<sup>42</sup>, at each occurrence, is independently selected from
                    H, CF<sub>3</sub>, halo, OH, CO<sub>2</sub>H, SO<sub>2</sub>R<sup>45</sup>, SOR<sup>45</sup>, SR<sup>45</sup>, NR<sup>46</sup>SO<sub>2</sub>R<sup>45</sup>.
                    NR<sup>46</sup>COR<sup>45</sup>, NR<sup>46</sup>R<sup>47</sup>, NO<sub>2</sub>, CN,
                    C<sub>1-4</sub> alkyl, C<sub>2-6</sub> alkenyl, C<sub>2-6</sub> alkynyl, C<sub>1-4</sub> alkoxy, and C<sub>1-4</sub> haloalkyl;
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 R^{45} is C_{1-4} alkyl;

R⁴⁶, at each occurrence, is independently selected from H and C₁₋₄ alkyl;

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R⁴⁷, at each occurrence, is independently selected from H, C₁₋₄ alkyl,

 $-C(=O)NH(C_{1-4} \text{ alkyl}), -SO_2(C_{1-4} \text{ alkyl}),$

-C(=O)O(C_{1-4} alkyl), -C(=O)(C_{1-4} alkyl), and -C(=O)H;

10 m is 1 or 2;

provided that when R^{11} is C_{1-6} alkyl, then R^{1} is not a C_{1-4} alkyl substituted by a) an unsubstituted 3H-pyrimidine-4-one moiety, b) a substituted 3H-pyrimidine-4-one moiety, c) an unsubstituted bicyclic derivative of 3H-pyrimidine-4-one, or d) a substituted bicyclic derivative of 3H-pyrimidine-4-one;

provided that when R^6 is -O-R¹¹ and R^6 is C₁₋₆ alkyl; then R^{8a} is not a substituted or unsubstituted indole moiety.

In another embodiment, the present invention provides a novel compound of Formula (Ia) wherein:

$$R^{8}$$
 R^{9}
 R^{1}
 R^{4a}
 R^{7}
 R^{6}
 R^{5}
 R^{1}
 R^{4a}

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or a stereoisomer or a pharmaceutically acceptable salt form thereof, wherein:

R¹ is selected from

H, C₁₋₃ haloalkyl, C₃₋₆ cycloalkyl,

C₁₋₄ alkyl substituted with 0-2 R²,

5 C₂₋₄ alkenyl substituted with 0-2 R², and

C₂₋₄ alkynyl substituted with 0-2 R²;

R², at each occurrence, is independently selected from

halo, C₁₋₃ haloalkyl, C₁₋₄ alkoxy, C₁₋₄ alkyl,

10 C₃₋₆ cycloalkyl, and phenyl substituted with 0-5 R⁴²;

R^{4a} is H or C₁₋₄ alkyl;

R⁵ is H, C₁₋₄ alkyl substituted with 0-2 R²⁰, or C₁₋₄ haloalkyl;

15

R⁶ is selected from

halo, -CF3, -OCF3, -CN, -NO2, -OCH3, -SCH3, -CF2CF3, -O-R11,

-OCF2CF3, -OCF2H, -OCF2CH3,

 $-S-R^{11}$, $-S(=O)-R^{11}$, $-S(=O)_2-R^{11}$, $-NR^{10}-R^{11}$,

20 -CH₂O-R¹¹, -CH₂S-R¹¹, CH₂S(=O)-R¹¹, CH₂S(=O)₂-R¹¹,

-CH2NR¹⁰-R¹¹,

C₁₋₄ haloalkyl, (C₁₋₄ haloalkyl)oxy;

C₁₋₄ alkyl substituted with 0-2 R²⁰,

C₂₋₄ alkenyl substituted with 0-2 R²⁰,

25 C₂₋₄ alkynyl substituted with 0-1 R²⁰, and

 C_{3-6} carbocyclic residue substituted with 0-3 R^{21} ,

 R^7 and R^9 are independently selected from

H, F, Cl, Br, -CF3, -OCF3, -OH, -CN, -NO2, -CF2CF3,C₁₋₄ alkyl, C₂₋₄ alkenyl, C₂₋₄ alkynyl, C₁₋₄ haloalkyl, C₁₋₄ alkoxy, and (C₁₋₄ haloalkyl)oxy;

5 R⁸ is selected from

halo, -CF3, -OCF3, -OH, -CN, -NO2, -OCH3, -SCH3, -CF2CF3,
-OR¹², -SR¹², -NR¹²R¹³, -C(O)H, -C(O)R¹², -C(O)NR¹²R¹³,
-NR¹⁴C(O)R¹², -C(O)OR¹², -OC(O)R¹², -OC(O)OR¹²,
-S(O)R¹², -S(O)2R¹², -S(O)NR¹²R¹³, -S(O)2NR¹²R¹³,
-NR¹⁴S(O)R¹², -NR¹⁴S(O)2R¹², -NR¹²C(O)R¹⁵, -NR¹²C(O)OR¹⁵,
-NR¹²S(O)2R¹⁵, -NR¹²C(O)NHR¹⁵;
C1-6 alkyl substituted with 0-2 R^{8a},
C2-6 alkenyl substituted with 0-2 R^{8a},
C2-6 alkynyl substituted with 0-2 R^{8a},
C3-6 cycloalkyl substituted with 0-2 R^{8a},
and
C3-10 carbocyclic residue substituted with 0-3 R³³;

R^{8a}, at each occurrence, is independently selected from halo, -CF₃, -OCF₃, -OH, -CN, -NO₂, -CF₂CF₃,

20 methyl, ethyl, n-propyl, i-propyl, n-butyl, i-butyl, s-butyl, t-butyl,
-OR¹², -SR¹², -NR¹²R¹³, -C(O)H, -C(O)R¹², -C(O)NR¹²R¹³,
-NR¹⁴C(O)R¹², -C(O)OR¹², -OC(O)R¹², -OC(O)OR¹²,
-S(O)R¹², -S(O)₂R¹², -S(O)NR¹²R¹³, -S(O)₂NR¹²R¹³,
-NR¹⁴S(O)R¹², -NR¹⁴S(O)₂R¹², -NR¹²C(O)R¹⁵, -NR¹²C(O)OR¹⁵,
-NR¹²S(O)₂R¹⁵, -NR¹²C(O)NHR¹⁵;

phenyl substituted with 0-5 R³³;

 C_{3-10} carbocyclic residue substituted with 0-3 R^{33} , and

5-6 membered heterocyclic ring system containing from 1-4 heteroatoms selected from the group consisting of N, O, and S substituted with 0-3 R³³;

5 R^{10} is H or C_{1-4} alkyl;

R¹¹ is selected from

C₁₋₆ alkyl substituted with 0-2 R²⁰,

C₂₋₆ alkenyl substituted with 0-2 R²⁰,

10 C₂₋₆ alkynyl substituted with 0-1 R²⁰,

C₃₋₁₀ carbocyclic residue substituted with 0-3 R²¹.

aryl substituted with 0-5 R²³, and

5-10 membered heterocyclic ring system containing from 1-4 heteroatoms selected from the group consisting of N, O, and S substituted with 0-3 R²¹:

·

alternatively, R^{10} and R^{11} join to form a 5- or 6-membered ring optionally substituted with -O- or -N(R^{14})-;

alternatively, R¹⁰ and R¹¹ when attached to N may be combined to form a 9- or 10membered bicyclic heterocyclic ring system containing from 1-3 heteroatoms
selected from the group consisting of N, O, and S, wherein said bicyclic
heterocyclic ring system is unsaturated or partially saturated, wherein said
bicyclic heterocyclic ring system is substituted with 0-3 R¹⁶;

25

15

R¹² is selected from H,

C₁₋₆ alkyl substituted with 0-2 R^{12a},

C₂₋₆ alkenyl substituted with 0-2 R^{12a},

C₂₋₆ alkynyl substituted with 0-2 R^{12a},

10

20

C₃₋₆ cycloalkyl substituted with 0-3 R³³, arvl substituted with 0-5 R³³;

C₃₋₁₀ carbocyclic residue substituted with 0-3 R³³, and

5-10 membered heterocyclic ring system containing from 1-4 heteroatoms selected from the group consisting of N, O, and S substituted with 0-3 R³³;

 R^{12a} , at each occurrence, is independently selected from

$$-SO_2NR^{46}R^{47}$$
, $-CONR^{46}R^{47}$, $-OR^{45}$, $=O$,

C₁₋₄ alkyl, C₂₋₆ alkenyl, C₂₋₆ alkynyl,

phenyl substituted with 0-5 R³³;

C₃₋₁₀ carbocyclic residue substituted with 0-3 R³³, and

5-10 membered heterocyclic ring system containing from 1-4 heteroatoms selected from the group consisting of N, O, and S substituted with 0-3 R³³;

R¹³, at each occurrence, is independently selected from H, C₁₋₄ alkyl, C₂₋₄ alkenyl, and C₂₋₄ alkynyl;

alternatively, R^{12} and R^{13} join to form a 5- or 6-membered ring optionally substituted with -O- or -N(R^{14})-;

alternatively, R¹² and R¹³ when attached to N may be combined to form a 9- or 10-membered bicyclic heterocyclic ring system containing from 1-3 heteroatoms selected from the group consisting of N, O, and S, wherein said bicyclic heterocyclic ring system is unsaturated or partially saturated, wherein said bicyclic heterocyclic ring system is substituted with 0-3 R¹⁶;

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R<sup>14</sup>, at each occurrence, is independently selected from H and C<sub>1-4</sub> alkyl;
R<sup>15</sup>, at each occurrence, is independently selected from
          H, C<sub>1-4</sub> alkyl, C<sub>2-4</sub> alkenyl, and C<sub>2-4</sub> alkynyl;
R<sup>16</sup>, at each occurrence, is independently selected from
          H, OH, halo, CN, NO<sub>2</sub>, CF<sub>3</sub>, SO<sub>2</sub>R<sup>45</sup>, NR<sup>46</sup>R<sup>47</sup>, -C(=O)H,
          C<sub>1-4</sub> alkyl, C<sub>2-4</sub> alkenyl, C<sub>2-4</sub> alkynyl, C<sub>1-4</sub> haloalkyl,
          C<sub>1-3</sub> haloalkyl-oxy-, and C<sub>1-3</sub> alkyloxy-;
R<sup>20</sup> is selected from
          H, halo, -OH, -CF3, -CN, -NO2, -CO2H, -SO2R<sup>45</sup>,
          -SOR45, -SR45, -NR46SO2R45, -NR46COR45, -NR46R47,
          C<sub>1-4</sub> alkyl, C<sub>2-6</sub> alkenyl, C<sub>2-6</sub> alkynyl, C<sub>1-4</sub> alkoxy,
          C<sub>1-4</sub> haloalkyl;
          C<sub>3-10</sub> carbocyclic residue substituted with 0-3 R<sup>21</sup>:
          aryl substituted with 0-5 R<sup>23</sup>; and
          5-10 membered heterocyclic ring system containing from 1-4 heteroatoms
                    selected from the group consisting of N, O, and S substituted with 0-3
                    R^{21}:
R^{21}, at each occurrence, is independently selected from
          H. OH. halo, CF<sub>3</sub>, SO<sub>2</sub>R<sup>45</sup>, NR<sup>46</sup>R<sup>47</sup>, CN, NO<sub>2</sub>, =O, C<sub>1-4</sub> alkyl;
          C<sub>1-4</sub> alkoxy, and (C<sub>1-4</sub> haloalkyl)oxy;
```

 R^{23} , at each occurrence, is independently selected from H, OH, halo, CF3, SO_2R^{45} , $NR^{46}R^{47}$, CN, NO_2 , C_{1-4} alkyl; C_{1-4} alkoxy, and $(C_{1-4}$ haloalkyl)oxy;

```
R<sup>33</sup>, at each occurrence, is independently selected from
                   H, OH, halo, -CN, -NO<sub>2</sub>, -CF<sub>3</sub>, -OCF<sub>3</sub>, -SO<sub>2</sub>R<sup>35</sup>, -S(=O)R<sup>35</sup>,
                   -SR^{35}, -NR^{36}R^{37}, -NHC(=O)R^{35}, -C(=O)NR^{36}R^{37}.
                   -C(=O)H, -C(=O)R^{35}, -C(=O)OR^{35}, -OC(=O)R^{35}, -OR^{35}.
  5
                   C<sub>1-6</sub> alkyl, C<sub>2-6</sub> alkenyl, C<sub>2-6</sub> alkynyl, C<sub>1-4</sub> haloalkyl,
                   C<sub>1-4</sub> alkoxy, (C<sub>1-4</sub> haloalkyl)oxy,
                   C<sub>3-6</sub> cycloalkyl, phenyl, aryl substituted with 0-2 R<sup>34</sup>.
                   C<sub>1-6</sub> alkyl substituted with R<sup>34</sup>, and
                   C<sub>2-6</sub> alkenyl substituted with R<sup>34</sup>;
10
        R<sup>34</sup>, at each occurrence, is independently selected from
                   OH, C<sub>1-4</sub> alkoxy, -SO<sub>2</sub>R<sup>35</sup>, -NR<sup>36</sup>R<sup>37</sup>, -NR<sup>36</sup>R<sup>37</sup>C(=O)-, and
                   (C<sub>1-4</sub> alkyl)CO<sub>2</sub>-;
15
        R<sup>35</sup>, at each occurrence, is independently selected from
                   C<sub>1-4</sub> alkyl, C<sub>1-4</sub> haloalkyl, C<sub>3-6</sub> cycloalkyl,
                   (C<sub>3-6</sub> cycloalkyl)methyl-, and (C<sub>3-6</sub> cycloalkyl)ethyl-;
        R<sup>36</sup>, at each occurrence, is independently selected from H and C<sub>1-4</sub> alkyl;
20
        R^{37}, at each occurrence, is independently selected from H, C_{1-4} alkyl,
                   -C(=O)NH(C_{1-4} \text{ alkyl}), -SO_2(C_{1-4} \text{ alkyl}), -C(=O)O(C_{1-4} \text{ alkyl}),
                   -C(=O)( C<sub>1-4</sub> alkyl), and -C(=O)H;
25
        R<sup>41</sup>, at each occurrence, is independently selected from
```

C₁₋₄ alkyl, C₂₋₈ alkenyl, C₂₋₈ alkynyl, C₁₋₄ alkoxy, and C₁₋₄ haloalkyl;

H, CF₃, halo, OH, CO₂H, SO₂R⁴⁵, NR⁴⁶R⁴⁷, NO₂, CN, =O.

R⁴², at each occurrence, is independently selected from H, CF₃, halo, OH, CO₂H, SO₂R⁴⁵, SOR⁴⁵, SR⁴⁵, NR⁴⁶SO₂R⁴⁵, NR⁴⁶COR⁴⁵, NR⁴⁶R⁴⁷, NO₂, CN, C₁₋₄ alkyl, C₂₋₆ alkenyl, C₂₋₆ alkynyl, C₁₋₄ alkoxy, and C₁₋₄ haloalkyl;

5

 R^{45} is C_{1-4} alkyl;

 R^{46} , at each occurrence, is independently selected from H and C_{1-4} alkyl;

10 R⁴⁷, at each occurrence, is independently selected from H, C₁₋₄ alkyl,

-C(=O)NH(C_{1-4} alkyl), -SO₂(C_{1-4} alkyl),

 $-C(=O)O(C_{1-4} \text{ alkyl}), -C(=O)(C_{1-4} \text{ alkyl}), \text{ and } -C(=O)H;$

m is 1 or 2.

15

In another embodiment, the present invention provides a novel compound of Formula (Ia) wherein:

$$R^{8}$$
 R^{7}
 R^{6}
 R^{5}
 R^{6}
 R^{5}
 R^{1}

20

or a stereoisomer or a pharmaceutically acceptable salt form thereof, wherein:

R¹ is selected from

25 H, CF3, methyl, ethyl, cyclopropyl, cyclobutyl, cyclopentyl, cyclohexyl,

C₁₋₄ alkyl substituted with 0-1 R²,

C₂₋₄ alkenyl substituted with 0-1 R², and

C₂₋₄ alkynyl substituted with 0-1 R²;

5 R² is selected from

F, Cl, CH₂F, CHF₂, CF₃, methyl, ethyl, cyclopropyl, cyclobutyl, cyclopentyl, cyclohexyl, and phenyl;

R^{4a} is H or methyl;

10

15

R⁵ is H, methyl, or ethyl;

R⁶ is selected from

 R^7 and R^9 are independently selected from

20 H, F, Cl, -CH₃, -OCH₃, -CF₃, -OCF₃, -CN, and -NO₂;

R⁸ is selected from

$$-OR^{12}$$
, $-SR^{12}$, $-NR^{12}R^{13}$, $-C(O)R^{12}$, $-S(O)R^{12}$, $-S(O)_2R^{12}$,

C₁₋₆ alkyl substituted with 0-2 R^{8a},

25 C₂₋₆ alkenyl substituted with 0-2 R^{8a},

C₂₋₆ alkynyl substituted with 0-2 R^{8a},

C₃₋₆ cycloalkyl substituted with 0-2 R^{8a}, and

C₃₋₁₀ carbocyclic residue substituted with 0-3 R³³;

R^{8a}, at each occurrence, is independently selected from halo, -CF₃, -OCF₃, -OH, -CN, -NO₂, -CF₂CF₃, methyl, ethyl, n-propyl, i-propyl, n-butyl, i-butyl, s-butyl, t-butyl, -OR¹², -SR¹², -NR¹²R¹³, -C(O)H, -C(O)R¹², -C(O)NR¹²R¹³, -NR¹⁴C(O)R¹², -C(O)OR¹², -OC(O)R¹², -OC(O)OR¹², -S(O)R¹², -S(O)₂R¹², -S(O)₂R¹², -S(O)₂R¹², -S(O)₂R¹², -NR¹²C(O)R¹³, -NR¹²C(O)R¹⁵, -NR¹²C(O)R¹⁵, -NR¹²C(O)R¹⁵, -NR¹²C(O)R¹⁵, -NR¹²C(O)R¹⁵, phenyl substituted with 0-5 R³³;

10 C₃₋₁₀ carbocyclic residue substituted with 0-3 R³³, and 5-6 membered heterocyclic ring system containing from 1-4 heteroatoms selected from the group consisting of N, O, and S substituted with 0-3 R³³;

15 R¹¹ is selected from

methyl, ethyl, propyl, and phenyl substituted with 0-5 R²³,

R¹² is selected from

C₁₋₆ alkyl substituted with 0-2 R^{12a},

20 C₂₋₆ alkenyl substituted with 0-2 R^{12a},

C₂₋₆ alkynyl substituted with 0-2 R^{12a},

C₃₋₆ cycloalkyl substituted with 0-3 R³³,

aryl substituted with 0-5 R³³;

C₃₋₁₀ carbocyclic residue substituted with 0-3 R³³, and

5-10 membered heterocyclic ring system containing from 1-4 heteroatoms selected from the group consisting of N, O, and S substituted with 0-3 R³³;

R^{12a}, at each occurrence, is independently selected from

H, halo, -OH, -CN, -NO₂, -CO₂H, -SO₂R⁴⁵, -SOR⁴⁵,
-SR⁴⁵, -NR⁴⁶SO₂R⁴⁵, -NR⁴⁶COR⁴⁵, -NR⁴⁶R⁴⁷,
-SO₂NR⁴⁶R⁴⁷, -CONR⁴⁶R⁴⁷, -OR⁴⁵, =O,
C₁₋₄ alkyl, C₂₋₆ alkenyl, C₂₋₆ alkynyl,

phenyl substituted with 0-5 R³³;
C₃₋₁₀ carbocyclic residue substituted with 0-3 R³³, and
5-10 membered heterocyclic ring system containing from 1-4 heteroatoms selected from the group consisting of N, O, and S substituted with 0-3 R³³;

10

15

 R^{13} , at each occurrence, is independently selected from H, C_{1-4} alkyl, C_{2-4} alkenyl, and C_{2-4} alkynyl;

alternatively, R¹² and R¹³ join to form a 5- or 6-membered ring selected from pyrrolyl, pyrrolidinyl, imidazolyl, piperidinyl, piperizinyl, methylpiperizinyl, and morpholinyl;

alternatively, R¹² and R¹³ when attached to N may be combined to form a 9- or 10membered bicyclic heterocyclic ring system containing from 1-3 heteroatoms
selected from the group consisting of N, O, and S; wherein said bicyclic
heterocyclic ring system is selected from indolyl, indolinyl, indazolyl,
benzimidazolyl, benzimidazolinyl, and benztriazolyl; wherein said bicyclic
heterocyclic ring system is substituted with 0-1 R¹⁶;

25 R¹⁴ is H, methyl, ethyl, propyl, or butyl;

R¹⁵ is H, methyl, ethyl, propyl, or butyl;

 R^{16} , at each occurrence, is independently selected from

H, OH, F, Cl, CN, NO₂, methyl, ethyl, methoxy, ethoxy, trifluoromethyl, and trifluoromethoxy;

 R^{23} , at each occurrence, is independently selected from

5 H, OH, F, Cl, CF₃, SO₂R⁴⁵, NR⁴⁶R⁴⁷, CN, NO₂, methyl, ethyl, propyl, and butyl;

 R^{33} , at each occurrence, is independently selected from

H, OH, halo, -CN, -NO₂, -CF₃, -OCF₃, -SO₂R³⁵, -S(=O)R³⁵,

 $-SR^{35}$, $-NR^{36}R^{37}$, $-NHC(=O)R^{35}$, $-C(=O)NR^{36}R^{37}$,

-C(=O)H, $-C(=O)R^{35}$, $-C(=O)OR^{35}$, $-OC(=O)R^{35}$, $-OR^{35}$,

C₁₋₆ alkyl, C₂₋₆ alkenyl, C₂₋₆ alkynyl, C₁₋₄ haloalkyl,

C₁₋₄ alkoxy, (C₁₋₄ haloalkyl)oxy,

C₃₋₆ cycloalkyl, phenyl, aryl substituted with 0-2 R³⁴,

15 C₁₋₆ alkyl substituted with R³⁴, and

C₂₋₆ alkenyl substituted with R³⁴;

R³⁴, at each occurrence, is independently selected from

OH,
$$C_{1-4}$$
 alkoxy, -SO₂R³⁵, -NR³⁶R³⁷, NR³⁶R³⁷C(=O)-, and (C_{1-4} alkyl)CO₂-;

 $\ensuremath{\mathsf{R}}^{35}\!,$ at each occurrence, is independently selected from

C₁₋₄ alkyl, C₁₋₄ haloalkyl, C₃₋₆ cycloalkyl,

(C3-6 cycloalkyl)methyl-, and (C3-6 cycloalkyl)ethyl-;

25

20

R³⁶, at each occurrence, is independently selected from H and C₁₋₄ alkyl;

 R^{37} , at each occurrence, is independently selected from H, C_{1-4} alkyl,

 $-C(=O)NH(C_{1-4} \text{ alkyl}), -SO_2(C_{1-4} \text{ alkyl}),$

 $-C(=O)O(C_{1-4} \text{ alkyl}), -C(=O)(C_{1-4} \text{ alkyl}), \text{ and } -C(=O)H;$

 R^{45} is C_{1-4} alkyl;

5 R⁴⁶, at each occurrence, is independently selected from H and C₁₋₄ alkyl;

R⁴⁷, at each occurrence, is independently selected from H, C₁₋₄ alkyl,

$$-C(=O)O(C_{1-4} \text{ alkyl}), -C(=O)(C_{1-4} \text{ alkyl}), \text{ and } -C(=O)H;$$

m is 1 or 2.

In another embodiment, the present invention provides a novel compound of Formula (Ib) wherein:

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$$R^8$$
 R^9
 R^7
 R^6
 R^5
 R^5
 R^6

or a stereoisomer or a pharmaceutically acceptable salt form thereof, wherein:

20

R¹ is selected from H, methyl, and ethyl;

R⁵ is H, methyl, or ethyl;

25 R⁶ is selected from

```
-NO<sub>2</sub>, -OCH<sub>3</sub>, -OCH<sub>2</sub>CH<sub>3</sub>, -SCH<sub>3</sub>, -SCH<sub>2</sub>CH<sub>3</sub>, -S(=O)CH<sub>3</sub>,
                   -S(=O)<sub>2</sub>CH<sub>3</sub>, methyl, ethyl, n-propyl, i-propyl, n-butyl, i-butyl, and s-butyl;
        R<sup>7</sup> is H. F. or Cl:
  5
        R<sup>8</sup> is selected from
                  -OR^{12}, -SR^{12}, -NR^{12}R^{13}, -C(O)R^{12}, -S(O)R^{12}, -S(O)2R^{12},
                  C<sub>1-6</sub> alkyl substituted with 0-2 R<sup>8a</sup>,
                  C<sub>3-6</sub> cycloalkyl substituted with 0-2 R<sup>8a</sup>, and
                  C<sub>3-10</sub> carbocyclic residue substituted with 0-3 R<sup>33</sup>;
10
        R<sup>8a</sup>, at each occurrence, is independently selected from
                  H, F, Cl, Br, methyl, ethyl, n-propyl, i-propyl, n-butyl, i-butyl, s-butyl,
                  t-butyl, -OH, methoxy, ethoxy, n-propoxy, i-propoxy, -CF3, -OCF3,
15
                  -CN, -NO<sub>2</sub>, -CF<sub>2</sub>CF<sub>3</sub>,-SCH<sub>3</sub>, -SCH<sub>2</sub>CH<sub>3</sub>, -SO<sub>2</sub>CH<sub>3</sub>, -NH<sub>2</sub>,
                  -CH2NH(CH3), -CH2N(CH3)2, -NH(CH3), -N(CH3)2, -CO(CH3),
                  -CO(OCH<sub>3</sub>), -NHCO(CH<sub>3</sub>), -CONH<sub>2</sub>, -C(=O)H, -CH(OH)CH<sub>3</sub>, -CH<sub>2</sub>OH,
                  -CH2CH2OH, -CH2OCH3, -CH2CH2OCH3, -CH2OCH2CH3,
                  phenyl substituted with 0-5 R<sup>33</sup> and pyridyl substituted with 0-5 R<sup>33</sup>
20
        R^9 is H;
        R<sup>12</sup> is selected from
                  C<sub>1-6</sub> alkyl substituted with 0-2 R<sup>12a</sup>,
                  cyclopropyl substituted with 0-2 R<sup>33</sup>.
25
                  cyclobutyl substituted with 0-2 R<sup>33</sup>,
                  cyclopentyl substituted with 0-2 R<sup>33</sup>,
                  cyclohexyl substituted with 0-2 R<sup>33</sup>,
                  bicyclo[3.1.1]heptane substituted with 0-2 R<sup>33</sup>,
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bicyclo[2.2.1]heptane substituted with 0-2 R³³, phenyl substituted with 0-3 R³³; and pyridyl substituted with 0-3 R³³;

- 5 R^{12a}, at each occurrence, is independently selected from H, F, Cl, -OH, methyl, ethyl, cyclopropyl substituted with 0-2 R³³, cyclobutyl substituted with 0-2 R³³, cyclopentyl substituted with 0-2 R³³, cyclohexyl substituted with 0-2 R³³, bicyclo[3.1.1]heptane substituted with 0-2 R³³, bicyclo[2.2.1]heptane substituted with 0-2 R³³, and phenyl substituted with 0-3 R³³:
- 15 R¹³ is H, methyl, or ethyl;

25

R³³, at each occurrence, is independently selected from
H, F, Cl, Br, methyl, ethyl, n-propyl, i-propyl,
n-butyl, i-butyl, s-butyl, t-butyl, -OH, methoxy, ethoxy, n-propoxy,
i-propoxy, -SCH₃, -SCH₂CH₃, -SO₂CH₃, -CF₃, -OCF₃, -CF₂CF₃, -CN,
-NO₂, -NH₂, -CH₂NH(CH₃), -CH₂N(CH₃)₂,
-NH(CH₃), -N(CH₃)₂, -CO(CH₃), -CO(OCH₃), -NHCO(CH₃),
-CONH₂, -C(=O)H, -CH(OH)CH₃, -CH₂OH, -CH₂CH₂OH,
-CH₂OCH₃, -CH₂CH₂OCH₃, and -CH₂OCH₂CH₃.

In another embodiment, the present invention provides a novel compound of Formula (Ib):

$$R^{8}$$
 R^{7}
 R^{6}
 R^{5}
 R^{5}
 R^{7}
 R^{1}

or a stereoisomer or a pharmaceutically acceptable salt form thereof, wherein:

5

R¹ is H or methyl;

R⁵ is H or methyl;

10 R⁶ is selected from

-F, -Cl, -CF₃, -CF₂CF₃, -OCF₃, -OCF₂CF₃, -OCF₂H, -OCF₂CH₃, -CN, -OCH₃, -SCH₃, -S(=O)CH₃, -S(=O)2CH₃, or methyl;

 R^7 is H, F, or Cl;

15

R⁸ is selected from

 $-OR^{12}$, $-SR^{12}$, $-NR^{12}R^{13}$,

 C_{1-6} alkyl substituted with 0-2 R^{8a} , and

 C_{3-6} cycloalkyl substituted with 0-2 R^{8a} ,

20

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 R^{8a} , at each occurrence, is independently selected from

H, F, Cl, Br, methyl, ethyl, n-propyl, i-propyl, n-butyl, i-butyl, s-butyl,

 $t\hbox{-butyl, -OH, methoxy, ethoxy, n-propoxy, i-propoxy, -CF3, -OCF3,}\\$

-CN, -CF2CF3,-SCH3, -SCH2CH3, -CH2NH(CH3), -CH2N(CH3)2,

-NH(CH3), -N(CH3)2, -CO(CH3), -CO(OCH3), -NHCO(CH3), -CONH2,

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-CH(OH)CH<sub>3</sub>, -CH<sub>2</sub>OH, -CH<sub>2</sub>CH<sub>2</sub>OH, -CH<sub>2</sub>OCH<sub>3</sub>, -CH<sub>2</sub>CCH<sub>2</sub>OCH<sub>3</sub>, -CH<sub>2</sub>OCH<sub>2</sub>CH<sub>3</sub>. phenyl substituted with 0-5 R<sup>33</sup>, and pyridyl substituted with 0-5 R<sup>33</sup>
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5 R^9 is H;

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R¹² is selected from
methyl, ethyl, propyl, butyl, pentyl, hexyl,
cyclopropyl, cyclobutyl, cyclopentyl, cyclohexyl,
bicyclo[3.1.1]heptane, bicyclo[2.2.1]heptane,
methyl substituted with R^{12a};
ethyl substituted with R^{12a};
propyl substituted with R^{12a};
phenyl substituted with 0-2 R³³; and
pyridyl substituted with 0-2 R³³;

R^{12a} is selected from cyclopropyl, cyclobutyl, cyclopentyl, cyclohexyl, bicyclo[3.1.1]heptane, bicyclo[2.2.1]heptane, and phenyl substituted with 0-2 R³³;

 R^{13} is H, methyl, or ethyl;

R³³, at each occurrence, is independently selected from

H, F, Cl, Br, methyl, ethyl, n-propyl, i-propyl, n-butyl, i-butyl, s-butyl,
t-butyl, -OH, methoxy, ethoxy, n-propoxy, i-propoxy, -SCH₃, -SCH₂CH₃,
-SO₂CH₃, -CF₃, -OCF₃, -CN, -NO₂, -NH₂, -CH₂NH(CH₃),
-CH₂N(CH₃)₂, -NH(CH₃), -N(CH₃)₂, -CO(CH₃), -CO(OCH₃),
-NHCO(CH₃), -CONH₂, -C(=O)H, -CH(OH)CH₃, -CH₂OH, -CH₂CH₂OH,
-CH₂OCH₃, -CH₂CH₂OCH₃, and -CH₂OCH₂CH₃.

In another embodiment, the present invention provides a novel compound of Formula (Ic):

$$R^{8}$$
 R^{9}
 R^{7}
 R^{1}
 R^{1}
 R^{1}
 R^{1}
 R^{1}
 R^{2}
 R^{3}
 R^{4}
 R^{5}
 R^{5}
(Ic)

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or a pharmaceutically acceptable salt thereof.

In an even further more preferred embodiment of the present invention, are compounds of Formula (I) selected from disclosed Examples 1-212.

In another embodiment R^1 is H or methyl.

In another embodiment R¹ is methyl.

In another embodiment R¹ is H.

In another embodiment R^5 is H or methyl.

In another embodiment R^5 is methyl.

In another embodiment R⁵ is H.

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In another embodiment R⁶ is -CF₃, -OCF₃, -CN, -OCH₃, -SCH₃, -S(=O)CH₃, -S(=O)₂CH₃, or methyl.

In another embodiment R⁶ is -CF₃.

In another embodiment R⁶ is -CN.

25 In another embodiment R⁶ is -OCH₃.

In another embodiment R⁶ is -SCH₃.

In another embodiment R⁶ is methyl.

In another embodiment R⁷ is H.

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In another embodiment R⁹ is H.

In another embodiment R^8 is -OR¹², -SR¹², -NR¹²R¹³, C_{1-6} alkyl substituted with 0-2 R^{8a} , or C_{3-6} cycloalkyl substituted with 0-2 R^{8a} .

In another embodiment R^8 is $-OR^{12}$.

In another embodiment R⁸ is -SR¹².

In another embodiment R^8 is $-NR^{12}R^{13}$.

In another embodiment R^8 is C_{1-6} alkyl substituted with 0-2 R^{8a} .

In another embodiment R⁸ is C₃₋₆ cycloalkyl substituted with 0-2 R^{8a}.

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In another embodiment R^{8a}, at each occurrence, is independently selected from H, F, Cl, Br, methyl, ethyl, n-propyl, i-propyl, n-butyl, i-butyl, s-butyl, t-butyl, -OH, methoxy, ethoxy, n-propoxy, i-propoxy, -CF3, -OCF3, -CN, -CF2CF3, -SCH3, -SCH2CH3, -CH2NH(CH3), -CH2N(CH3)2, -NH(CH3), -N(CH3)2, -CO(CH3), -CO(OCH3), -NHCO(CH3), -CONH2, -CH(OH)CH3, -CH2OH, -CH2CH2OH, -CH2OCH3, -CH2CH2OCH3, -CH2OCH2CH3, phenyl substituted with 0-5 R³³, and pyridyl substituted with 0-5 R³³

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In another embodiment R^{12} is selected from methyl substituted with R^{12a} ; ethyl substituted with R^{12a} ; propyl substituted with R^{12a} ; and phenyl substituted with R^{12a} .

In another embodiment R^{12} is phenyl substituted with 0-3 R^{33} .

In another embodiment R^{12} is phenyl substituted with 0-2 R^{33} .

In another embodiment R³³, at each occurrence, is independently selected from H, F, Cl, Br, methyl, ethyl, n-propyl, i-propyl, n-butyl, i-butyl, s-butyl, t-butyl, -OH, methoxy, ethoxy, n-propoxy, i-propoxy, -SCH3, -SCH2CH3, -SO2CH3, -SO2CH2CH3, -CF3, -OCF3, -CN, -NO2, -NH2, -CH2NH(CH3), -CH2N(CH3)2, -NH(CH3), -N(CH3)2, -CO(CH3), -CO(OCH3), -NHCO(CH3), -CONH2, -C(=O)H, -CH(OH)CH3, -CH2OH,

-CH2CH2OH, -CH2OCH3, -CH2CH2OCH3, and -CH2OCH2CH3.

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In a second embodiment, the present invention provides a pharmaceutical composition comprising a compound of Formula (I) and a pharmaceutically acceptable carrier.

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In a third embodiment, the present invention provides a method for the treatment a central nervous system disorder comprising administering to a host in need of such treatment a therapeutically effective amount of a compound of Formula (I), or a pharmaceutically acceptable salt thereof, wherein the compound is a 5HT_{2C} agonist.

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In a more preferred embodiment the present invention provides a method for the treatment central nervous system disorders including obesity, anorexia, bulemia, depression, anxiety, psychosis, schizophrenia, migraine, addictive behavior, obsessive-compulsive disorder, and sexual disorders comprising administering to a host in need of such treatment a therapeutically effective amount of a compound of Formula (I).

comprises obesity.

comprises schizophrenia.

In another further preferred embodiment the central nervous system disorder

In a further preferred embodiment the central nervous system disorder

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In another further preferred embodiment the central nervous system disorder comprises depression.

In another further preferred embodiment the central nervous system disorder comprises anxiety.

In another further preferred embodiment the central nervous system disorder sexual disorders.

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In another further preferred embodiment the central nervous system disorder addictive behaviors.

In a fourth embodiment the present invention provides novel compounds of Formula (I) or pharmaceutically acceptable salt forms thereof for use in therapy.

In a fifth embodiment the present invention provides the use of novel compounds of Formula (I) or pharmaceutically acceptable salt forms thereof for the manufacture of a medicament for the treatment of central nervous system disorders including obesity, anorexia, bulemia, depression, anxiety, psychosis, schizophrenia, migraine, addictive behavior, obsessive-compulsive disorder, and sexual disorders.

DEFINITIONS

25 of the present invention containing an asymmetrically substituted atom may be isolated in optically active or racemic forms. It is well known in the art how to prepare optically active forms, such as by resolution of racemic forms or by synthesis from optically active starting materials. Many geometric isomers of olefins, C=N double bonds, and the like can also be present in the compounds described herein, and all such stable isomers are contemplated in the present invention. *Cis* and *trans* geometric isomers of the compounds of the present invention are described and may be isolated as a mixture of isomers or as separated isomeric forms. All chiral,

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diastereomeric, racemic forms and all geometric isomeric forms of a structure are intended, unless the specific stereochemistry or isomeric form is specifically indicated.

The term "substituted", as used herein, means that any one or more hydrogens on the designated atom is replaced with a selection from the indicated group, provided that the designated atom's normal valency is not exceeded, and that the substitution results in a stable compound. When a substituent is keto (i.e., =O), then 2 hydrogens on the atom are replaced.

When any variable (e.g. R², R²³, R³³ etc.) occurs more than one time in any constituent or formula for a compound, its definition at each occurrence is independent of its definition at every other occurrence. Thus, for example, if a group is shown to be substituted with 0-2 R², then said group may optionally be substituted zero, one or two R² groups and R² at each occurrence is selected independently from the definition of R². Also, combinations of substituents and/or variables are permissible only if such combinations result in stable compounds.

When a bond to a substituent is shown to cross a bond connecting two atoms in a ring, then such substituent may be bonded to any atom on the ring. When a substituent is listed without indicating the atom via which such substituent is bonded to the rest of the compound of a given formula, then such substituent may be bonded via any atom in such substituent. Combinations of substituents and/or variables are permissible only if such combinations result in stable compounds.

As used herein, "alkyl" or "alkylene" is intended to include both branched and straight-chain saturated aliphatic hydrocarbon groups having the specified number of carbon atoms; for example, "C1-C6 alkyl" or "C1-6 alkyl" denotes alkyl having 1 to 6 carbon atoms. Examples of alkyl include, but are not limited to, methyl, ethyl, n-propyl, i-propyl, n-butyl, i-butyl, sec-butyl, t-butyl, n-pentyl, n-hexyl, 2-methylbutyl, 2-methylpentyl, 2-ethylbutyl, 3-methylpentyl, and 4-methylpentyl. Unless otherwise specified, preferred alkyl are methyl, ethyl, n-propyl, i-propyl, n-butyl, i-butyl, sec-butyl, and t-butyl.

"Alkenyl" or "alkenylene" is intended to include hydrocarbon chains of either a straight or branched configuration having the specified number of carbon atoms, for example "C2-6 alkenyl", and one or more unsaturated carbon-carbon bonds which

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may occur in any stable point along the chain. Examples of alkenyl include, but are not limited to, ethenyl, 1-propenyl, 2-propenyl, 2-butenyl, 3-butenyl, 2-pentenyl, 3, pentenyl, 4-pentenyl, 2-hexenyl, 3-hexenyl, 4-hexenyl, 5-hexenyl, 2-methyl-2-propenyl, 4-methyl-3-pentenyl, and the like. Unless otherwise specified, preferred alkenyl are ethenyl, 1-propenyl, 2-propenyl, 2-butenyl, and 3-butenyl.

"Alkynyl" or "alkynylene" is intended to include hydrocarbon chains of either a straight or branched configuration, having the specified number of carbon atoms, for example "C2-6 alkynyl", and one or more carbon-carbon triple bonds which may occur in any stable point along the chain, such as ethynyl, propynyl, butynyl, pentynyl, hexynyl and the like. Unless otherwise specified, preferred alkynyl are ethynyl and propynyl."

"Cycloalkyl" is intended to include saturated ring groups, having the specified number of carbon atoms. For example, "C3-C6 cycloalkyl" denotes such as cyclopropyl, cyclobutyl, cyclopentyl, or cyclohexyl.

"Alkoxy" or "alkyloxy" represents an alkyl group as defined above with the indicated number of carbon atoms attached through an oxygen bridge. Examples of alkoxy include, but are not limited to, methoxy, ethoxy, n-propoxy, i-propoxy, n-butoxy, s-butoxy, t-butoxy, n-pentoxy, and s-pentoxy. Unless otherwise specified, preferred alkoxy are methoxy, ethoxy, n-propoxy, and i-propoxy.

Similarly, "alkylthio" is represents an alkyl group as defined above with the indicated number of carbon atoms attached through a sulpher bridge. Unless otherwise specified, preferred alkylthio are methylthio and ethylthio.

"Halo" or "halogen" as used herein refers to fluoro, chloro, bromo, and iodo. Preferred halo are fluoro and chloro.

"Counterion" is used to represent a small, negatively charged species such as chloride, bromide, hydroxide, acetate, sulfate, and the like.

"Haloalkyl" is intended to include both branched and straight-chain saturated aliphatic hydrocarbon groups having the specified number of carbon atoms, substituted with 1 or more halogen (for example $-C_VF_W$ where v=1 to 3 and w=1 to (2v+1)). Examples of haloalkyl include, but are not limited to, trifluoromethyl, difluoromethyl, fluoromethyl, trichloromethyl, pentafluoroethyl, pentachloroethyl,

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2,2,2-trifluoroethyl, heptafluoropropyl, and heptachloropropyl. Unless otherwise specified, preferred haloalkyl are trifluoromethyl, difluoromethyl, and fluoromethyl.

As used herein, "carbocycle" or "carbocyclic residue" or "carbocyclic moiety" is intended to mean any stable 3- to 7-membered monocyclic or bicyclic or 7- to 13-membered bicyclic or tricyclic, any of which may be saturated or partially unsaturated. Examples of such carbocycles include, but are not limited to, cyclopropyl, cyclobutyl, cyclopentyl, cyclohexyl, cycloheptyl, cyclooctyl, cyclopentenyl, cyclohexenyl, bicyclo[3.1.1.]heptane, bicyclo[2.2.1]heptane, [3.3.0]bicyclooctane, [4.3.0]bicyclononane, [4.4.0]bicyclodecane (decalin), [2.2.2]bicyclooctane, indanyl, adamantyl, or tetrahydronaphthyl (tetralin).

As used herein, the term "heterocycle" or "heterocyclic ring" or "heterocyclic ring system" is intended to mean a stable 5- to 7- membered monocyclic or bicyclic or 7- to 14-membered bicyclic heterocyclic ring which is saturated, partially unsaturated, or unsaturated, and which consists of carbon atoms and 1, 2, 3 or 4 heteroatoms independently selected from the group consisting of N, O and S and including any bicyclic group in which any of the above-defined heterocyclic rings is fused to a benzene ring. The nitrogen and sulfur heteroatoms may optionally be oxidized. The heterocyclic ring may be attached to its pendant group at any heteroatom or carbon atom which results in a stable structure. The heterocyclic rings described herein may be substituted on carbon or on a nitrogen atom if the resulting compound is stable. If specifically noted, a nitrogen in the heterocycle may optionally be quaternized. It is preferred that when the total number of S and O atoms in the heterocycle exceeds one, then these heteroatoms are not adjacent to one another. It is preferred that the total number of S and O atoms in the heterocycle is not more than one.

Examples of heterocycles include, but are not limited to, 1H-indazole, 2-pyrrolidonyl, 2H,6H-1,5,2-dithiazinyl, 2H-pyrrolyl, 3H-indolyl, 4-piperidonyl, 4aH-carbazole, 4H-quinolizinyl, 6H-1,2,5-thiadiazinyl, acridinyl, azocinyl, benzimidazolyl, benzofuranyl, benzothiofuranyl, benzothiophenyl, benzoxazolyl, benzoxazolyl, benztriazolyl, benztriazolyl, benzisoxazolyl, benzisothiazolyl, benzimidazalonyl, carbazolyl, 4aH-carbazolyl, b-carbolinyl, chromanyl, chromenyl, cinnolinyl, decahydroquinolinyl, 2H,6H-1,5,2-dithiazinyl, dihydrofuro[2,3-b]tetrahydrofuran, furanyl, furazanyl, imidazolidinyl, imidazolinyl,

imidazolyl, imidazolopyridinyl, 1H-indazolyl, indolenyl, indolinyl, indolizinyl, indolyl, isatinoyl, isobenzofuranyl, isochromanyl, isoindazolyl, isoindolinyl, isoindolyl, isoquinolinyl, isothiazolyl, isothiazolopyridinyl, isoxazolyl, isoxazolopyridinyl, morpholinyl, naphthyridinyl, octahydroisoquinolinyl, oxadiazolyl, 5 1,2,3-oxadiazolyl, 1,2,4-oxadiazolyl, 1,2,5-oxadiazolyl, 1,3,4-oxadiazolyl, oxazolidinyl, oxazolyl, oxazolopyridinyl, oxazolidinylperimidinyl, oxindolyl, phenanthridinyl, phenanthrolinyl, phenarsazinyl, phenazinyl, phenothiazinyl, phenoxathiinyl, phenoxazinyl, phthalazinyl, piperazinyl, piperidinyl, pteridinyl, piperidonyl, 4-piperidonyl, pteridinyl, purinyl, pyranyl, pyrazinyl, pyrazolidinyl, 10 pyrazolinyl, pyrazolopyridinyl, pyrazolyl, pyridazinyl, pyridooxazole, pyridoimidazole, pyridothiazole, pyridinyl, pyridyl, pyrimidinyl, pyrrolidinyl, pyrrolinyl, pyrrolyl, quinazolinyl, quinolinyl, 4H-quinolizinyl, quinoxalinyl, quinuclidinyl, carbolinyl, tetrahydrofuranyl, tetrahydroisoguinolinyl, tetrahydroquinolinyl, 6H-1,2,5-thiadiazinyl, 1,2,3-thiadiazolyl, 1,2,4-thiadiazolyl, 15 1,2,5-thiadiazolyl, 1,3,4-thiadiazolyl, thianthrenyl, thiazolyl, thiazolopyridinyl, thienyl, thienothiazolyl, thienooxazolyl, thienoimidazolyl, thiophenyl, triazinyl, 1,2,3triazolyl, 1,2,4-triazolyl, 1,2,5-triazolyl, 1,3,4-triazolyl, and xanthenyl. Preferred 5 to 10 membered heterocycles include, but are not limited to, pyridinyl, pyrimidinyl, triazinyl, furanyl, thienyl, thiazolyl, pyrrolyl, pyrazolyl, pyrazinyl, piperazinyl, 20 imidazolyl, oxazolyl, isoxazolyl, tetrazolyl, benzofuranyl, benzothiofuranyl, indolyl, benzimidazolyl, 1*H*-indazolyl, oxazolidinyl, isoxazolidinyl, benzotriazolyl, benzisoxazolyl, oxindolyl, benzoxazolinyl, benzoxazolyl, benzthiazolyl, benzisothiazolyl, isatinoyl, isoxazolopyridinyl, isothiazolopyridinyl, thiazolopyridinyl, oxazolopyridinyl, imidazolopyridinyl, pyrazolopyridinyl, 25 quinolinyl, and isoquinolinyl. Preferred 5 to 6 membered heterocycles include, but are not limited to, pyridinyl, pyrimidinyl, triazinyl, furanyl, thienyl, thiazolyl, pyrrolyl, pyrazinyl, piperazinyl, piperidinyl, pyrazolyl, imidazolyl, oxazolyl, isoxazolyl, tetrazolyl, and oxazolidinyl; more preferred 5 to 6 membered heterocycles include, but are not limited to, pyridinyl, pyrimidinyl, triazinyl, furanyl, thienyl, 30 thiazolyl, piperazinyl, piperidinyl, pyrazolyl, imidazolyl, and tetrazolyl. Also included are fused ring and spiro compounds containing, for example, the above heterocycles.

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As used herein, the term "aryl", or aromatic residue, is intended to mean an aromatic moiety containing six to ten carbon atoms, such as phenyl and naphthyl.

The phrase "pharmaceutically acceptable" is employed herein to refer to those compounds, materials, compositions, and/or dosage forms which are, within the scope of sound medical judgment, suitable for use in contact with the tissues of human beings and animals without excessive toxicity, irritation, allergic response, or other problem or complication, commensurate with a reasonable benefit/risk ratio.

As used herein, "pharmaceutically acceptable salts" refer to derivatives of the disclosed compounds wherein the parent compound is modified by making acid or base salts thereof. Examples of pharmaceutically acceptable salts include, but are not limited to, mineral or organic acid salts of basic residues such as amines; alkali or organic salts of acidic residues such as carboxylic acids; and the like. The pharmaceutically acceptable salts include the conventional non-toxic salts or the quaternary ammonium salts of the parent compound formed, for example, from non-toxic inorganic or organic acids. For example, such conventional non-toxic salts include those derived from inorganic acids such as hydrochloric, hydrobromic, sulfuric, sulfamic, phosphoric, nitric and the like; and the salts prepared from organic acids such as acetic, propionic, succinic, glycolic, stearic, lactic, malic, tartaric, citric, ascorbic, pamoic, maleic, hydroxymaleic, phenylacetic, glutamic, benzoic, salicylic, sulfanilic, 2-acetoxybenzoic, fumaric, toluenesulfonic, methanesulfonic, ethane disulfonic, oxalic, isethionic, and the like.

The pharmaceutically acceptable salts of the present invention can be synthesized from the parent compound which contains a basic or acidic moiety by conventional chemical methods. Generally, such salts can be prepared by reacting the free acid or base forms of these compounds with a stoichiometric amount of the appropriate base or acid in water or in an organic solvent, or in a mixture of the two; generally, nonaqueous media like ether, ethyl acetate, ethanol, isopropanol, or acetonitrile are preferred. Lists of suitable salts are found in *Remington's Pharmaceutical Sciences*, 17th ed., Mack Publishing Company, Easton, PA, 1985, p. 1418, the disclosure of which is hereby incorporated by reference.

"Prodrugs" are intended to include any covalently bonded carriers which release the active parent drug according to Formula (I) *in vivo* when such prodrug is

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administered to a mammalian subject. Prodrugs of a compound of Formula (I) are prepared by modifying functional groups present in the compound in such a way that the modifications are cleaved, either in routine manipulation or *in vivo*, to the parent compound. Prodrugs include compounds of Formula (I) wherein a hydroxy, amino, or sulfhydryl group is bonded to any group that, when the prodrug or compound of Formula (I) is administered to a mammalian subject, cleaves to form a free hydroxyl, free amino, or free sulfhydryl group, respectively. Examples of prodrugs include, but are not limited to, acetate, formate and benzoate derivatives of alcohol and amine functional groups in the compounds of Formula (I), and the like.

"Stable compound" and "stable structure" are meant to indicate a compound that is sufficiently robust to survive isolation to a useful degree of purity from a reaction mixture, and formulation into an efficacious therapeutic agent.

SYNTHESIS

Throughout the details of the invention, the following abbreviations are used with the following meanings:

Reagents:

DIBAL diisobutyl aluminum hydride

20 Et₃N triethylamine

TFA trifluoroacetic acid

LAH lithium aluminum hydride

NBS N-bromosuccinimide

Red-Al Sodium bis(2-methoxyethoxy)aluminum hydride

25 Pd2dba3 Tris(dibenzylideneacetone)dipalladium(0)

ACE-Cl 2-chloroethylchloroformate

Solvents:

THF tetrahydrofuran

30 MeOH methanol

EtOH ethanol

EtOAc ethyl acetate

PH7483 NP

HOAc acetic acid

DMF dimethyl formamide

DMSO dimethyl sulfoxide

DME dimethoxyethane

5 Et₂O diethylether

iPrOH isopropanol

Others:

Ar aryl

10 Ph phenyl

Me methyl Et ethyl

NMR nuclear magnetic resonance

MHz megahertz

15 BOC tert-butoxycarbonyl

CBZ benzyloxycarbonyl

Bn benzyl
Bu butyl

Pr propyl

20 cat. catalytic

mL milliliter

nM nanometer

ppm part per million

mmol millimole

25 mg milligram

g gram

kg kilogram

TLC thin layer chromatography

HPLC high pressure liquid chromatography

30 rt room temperature

aq. aqueoussat. saturated

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The compounds of the present invention can be prepared in a number of ways well known to one skilled in the art of organic synthesis. The compounds of the present invention can be synthesized using the methods described below, together with synthetic methods known in the art of synthetic organic chemistry, or variations thereon as appreciated by those skilled in the art. Preferred methods include, but are not limited to, those described below. All references cited herein are hereby incorporated in their entirety herein by reference.

The novel compounds of this invention may be prepared using the reactions and techniques described in this section. The reactions are performed in solvents appropriate to the reagent and materials employed and are suitable for the transformations being effected. Also, in the description of the synthetic methods described below, it is to be understood that all proposed reaction conditions, including choice of solvent, reaction atmosphere, reaction temperature, duration of the experiment and workup procedures, are chosen to be the conditions standard for that reaction, which should be readily recognized by one skilled in the art. It is understood by one skilled in the art of organic synthesis that the functionality present on various portions of the molecule must be compatible with the regents and reactions proposed. Such restrictions to the substituents which are compatible with the reaction conditions will be readily apparent to one skilled in the art and alternate methods must then be used.

The preparation of compounds of Formula (I) of the present invention may be carried out in a convergent or sequential synthetic manner. Detailed synthetic preparations of the compounds of Formula (I) are shown in the following reaction schemes. The skills required in preparation and purification of the compounds of Formula (I) and the intermediates leading to these compounds are known to those in the art. Purification procedures include, but are not limited to, normal or reverse phase chromatography, crystallization, and distillation.

Several methods for the preparation of the compounds of the present invention are illustrated in the schemes and examples shown below. The substitutions are as described and defined above.

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Compounds of Formula (I) of this invention may be prepared as shown in Scheme 1. Thus, preparation of an aryl hydrazine (II) is accomplished, for example, by treatment of a corresponding substituted aniline with NaNO2 followed by reduction of the N-nitroso intermediate with SnCl₂ in conc. HCl. Assembly of the core indole intermediate (IV) is accomplished by Fischer indole cyclization of the aryl hydrazine and a suitably substituted ketone (i.e. (III)) by methods described by, but not limited to, R.J. Sundberg, "Indoles, Best Synthetic Methods" 1996, Academic Press, San Diego, CA. For example, treatment of the aryl hydrazine (II) as the free base or the corresponding mineral acid salt with the ketone (III) $(R^1 = H, Bn, CBZ)$. CO₂Et, etc) in an alcoholic solvent in the presence of mineral acid affords the indoles (IV) as the free bases (after treatment with aq. NaOH). Reduction of the indoles to the corresponding cis or trans substituted indolines is accomplished by, for example, treatment with hydrogen in the presence of a catalyst such as platinum oxide or palladium on carbon, or with a metal such as zinc and a mineral acid such as hydrochloric acid, or with sodium and liquid ammonia, or with borane-amine complex such as borane-triethylamine in tetrahydrofuran, or preferably by treatment with triethylsilane or NaCNBH3 in an acid such as acetic or trifluoroacetic acid.

SCHEME 1

R8 R9
$$R^9$$
 R^{4a} R^{4a}

N H⁵

(I)

The corresponding enantiomers can be isolated by separation of the racemic mixture of (I) on a chiral stationary phase column utilizing normal or reverse phase HPLC techniques, the details of which are described in the examples. Alternatively, a diastereomeric mixture of (I) can be prepared by treatment of (I, R² = H) with an appropriate chiral acid (or suitably activated derivative), for example dibenzoyl tartrate or the like (see, for example, Kinbara, K., et. al., *J. Chem. Soc., Perkin Trans.* 2, 1996, 2615; and Tomori, H., et. al., *Bull. Chem. Soc. Jpn.*, 1996, 3581). The diastereomers would then be separated by traditional techniques (i.e. silica chromatography, crystallization, HPLC, etc) followed by removal of the chiral

or base

2) R¹Cl, K₂CO₃, Kl, DMF

N R⁵

(V)

auxiliary to afford enantiomerically pure (I).

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In the cases where the carboline nitrogen has been protected (V) (i.e. Pg = Boc, Bn, CBZ, CO₂R), it may be removed under a variety of conditions as described

in Greene, T.W., Wuts, P.G.W., "Protective Groups in Organic Synthesis, 2nd Edition", John Wiley and Sons, Inc., New York, pages 309-405, **1991**. The free secondary amine could then be alkylated, for example, by treatment with a suitably substituted alkyl halide (R¹Cl, R¹Br or R¹I) and a base to afford additional compounds of type (I), as described, for example, by Glennon, R.A., et. al., *Med. Chem. Res.*, **1996**, 197.

Compounds of Formula (II) can be prepared as described in Scheme 2. Formation of the aryl amine (VII) may be accomplished by reduction of the corresponding aryl nitro compound (VI). The reduction may be accomplished with a variety of reducing agents, for example, LAH, SnCl₂, NaBH₄, N₂H₄, etc. or with hydrogen in the presence of a suitable catalyst, such as palladium on carbon, or platinum oxide, etc., (see Hudlicky, M., "Reductions in Organic Chemistry", Ellis Horwood, Ltd., Chichester, UK, 1984). Formation of the aryl hydrazine (II) may then be performed as previously described in Scheme 1 or more directly by treatment of the aniline (VII) with aq. hydrochloric acid, stannous chloride and NaNO₂ at room temperature (see, Buck, J.S., Ide, W.S., *Org. Syn., Coll. Vol.*, 2, 1943, 130). This latter procedure is especially important when initiating the synthesis with halogenated arylamines (VII). The necessity for preparation of the hydrazine intermediate without the use of strong reductive conditions is critical in these such examples.

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SCHEME 2

Another related route to hydrazines of Formula (II) is shown in Scheme 3. When an aromatic substitution pattern containing a sulfur moiety is desired the following route may be employed. Displacement of a halogen (Cl, F) of a suitably substituted aryl nitro derivative (VIII) by the prerequisite nucleophile under basic

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conditions affords intermediates of type (IX). Reduction of the nitro moiety followed by elaboration of the resultant amine to the substituted or unsubstituted hydrazine (X) is as described above.

5 SCHEME 3

Initiating the synthesis with a nitrobenzene derivative such as (VIII), this approach allows for a variety of derivatization. More highly substituted nitrobenzenes can be obtained by traditional synthetic manipulation (*i.e* aromatic substitution) and are known by those in the art (see Larock, R.C., Comprehensive Organic Transformations, VCH Publishers, New York, 1989).

An alternate, more direct approach to differentially substituted analogs is shown in Scheme 4. Initiating the preparation of compounds of type (I) with an aryl iodide expands the versatility of this approach. The preparation of an intermediate which can be functionalized at a later stage is a more efficient approach to some of the substitution types. Fischer indole cyclization of the iodide (XI) with the ketone (III) as described previously, followed by protection of the amine with Boc₂O, affords the iodo indole (XII). Alkylation of the indole nitrogen under basic conditions followed by removal of the Boc protecting group and a second alkylation of the carboline nitrogen affords the selective differentially substituted carboline indoles (XIII). Usual reduction of the indole to indoline is carried out without any loss of the aromatic halogen to afford the common aryliodide (XIV). Facile displacement of the iodide with a variety of sulfur nucleophiles under copper catalyzed conditions affords the diaryl sulfides (XV).

SCHEME 4

Additional method of preparing differentially substituted analogs is shown in Scheme 5. Facile displacement of the iodo-group of indole (XII) with Zn(CN)2 under Pd(0) catalyzed conditions affords the cyanoindole (XVI). Reduction of indole is achieved as described previously, followed by protection of the amine with Boc₂O and alkylation of the indoline nitrogen under basic conditions affords the cyanoindoline (XVIII). Alternatively, iodoindole (XII) can be reduced and treated Boc₂O to give iodoindoline (XVII) as described previously. Displacement of the iodide with Zn(CN)₂ catalyzed by Pd(0) under microwave condition followed by alkylation of the indoline nitrogen produces cyanoindoline (XVIII).

SCHEME 5

The preparation of compounds of Formula (I) with additional diversity of functionalization of the aromatic A ring of the tricycle is described here. Bromination of the indolines (I, R⁸ = H) when the amine is protected (with the Boc or CBZ protecting groups), with for example, NBS in DMF, affords the R⁸ brominated derivatives (XIX). These activated aryl derivatives (XIX) act as excellent counterparts for a number of important synthetic transformations.

One example is described in Scheme 6, where the aromatic ring of Formula (I) is substituted with an arylamino group. Treatment of bromide (XIX) with a variety of anilines (XX) in the presence of a Pd(0) catalyst, such as Pd₂(dba)₃, Pd(PPh₃)₄ or Pd(PPh₃)₂Cl₂, and suitable ligand such BINAP or PPh₃, and a base such as NaOtBu or CsCO₃ in a suitable solvent such as DMF, toluene, THF, DME, or the like, affords the biaryl anilines (XXI).

SCHEME 6

An alternate method for preparing biaryl anilines (XXI) is described in Scheme 7 and proceeds from brominated derivatives (XIX). Treatment of arylbromide derivatives of type (XIX) with diphenylmethyl imine in the presence of a Pd(0) catalyst, such as Pd2(dba)3, Pd(PPh3)4 or Pd(PPh3)2Cl2, and suitable ligand such BINAP or PPh3, and a base such as NaOtBu or CsCO3 in a suitable solvent such as DMF, toluene, THF, DME, or the like, followed affords an imine intermediate. Basic hydrolysis (hydroxylamine, sodium acetate in methanol) affords the primary aniline derivative (XXII). Coupling of these anilines with various arylbromide (XXIII) under Pd(0) catalyzed condition described above affords the biaryl anilines (XXI).

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SCHEME 7

Similarly arylamino coupling of the bromine derivatives (XXIV), readily obtained by the synthetic sequence exemplified in Scheme 2, (starting with the suitably functionalized bromo nitrobenzenes (VI), is shown in Scheme 8. This approach allows for the preparation of arylamino-indoles as well as the corresponding indoline derivatives. Protection of the amine functionality must be carried out if R¹, = H (see Greene et. al for protections of amines). This is readily accomplished, for example, by treatment of bromo derivatives (XXIV) with excess (Boc)₂O in aqueous sodium hydroxide and dioxane. Subsequent coupling with a variety of aryl anilines is carried out as described above in Scheme 6, to afford the arylamino indoline adducts (XXV). This protocol is amenable to R^a, R^b, and R^c bromide, iodide, triflates, and/or diazo derivatives. In addition, there exists a wide range of procedures and protocols for functionalizing haloaromatics, aryldiazonium and aryltriflate compounds. These procedures are well known by those in the art and described, for example (see Stanforth, S.P., Tetrahedron, 1998, 263; Buchwald, S.L., et. al., J. Am. Chem. Soc., 1998, 9722; Stille, J.K., et. al., J. Am. Chem. Soc., 1984, 7500). Among these procedures are biaryl couplings, alkylations, acylations, aminations, and amidations. The power of palladium catalyzed functionalization of aromatic cores has been explored in depth in the last decade. An excellent review of this field can be found in

J. Tsuji, "Palladium Reagents and Catalysts, Innovations in Organic Synthesis", J. Wiley and Sons, New York, 1995.

SCHEME 8

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The aniline (XXII) can also react with an appropriate aldehyde (XXVII) in the presence a suitable reducing agent such as sodium triacetoxyborohydride or sodium cyanoborohydride under mild reaction conditions, such as in the presence of acetic acid, in a suitable solvent such as 1,2-dichloroethane, THF, methanol or acetonitrile to produce the variety of secondary aniline analogs (XXVI). In analogy of Scheme 8, the protocol described in Scheme 9 can also be applied to analogs of (XIX) where the R⁷ or R⁹ groups are NH₂, Br, I, OTf, etc., to afford analogs of (XXVI) where the alkylamino group is on the R⁷ or R⁹ position.

SCHEME 9

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The aniline (XXII) can react with 1 equivalent of various alkylhalides or alkylsulfonates (XXVII) in the presence a suitable base such as NaH, K2CO3,

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Na₂CO₃, CsCO₃, Et₃N or Et₂(i-Pr)N in a suitable solvent such as DMF, DMSO, toluene, THF, DME or the like, produce the variety of secondary aniline analogs (XXVIII) as shown in scheme 10. In analogy of Scheme 8, the protocol described in Scheme 10 can also be applied to analogs of (XIX) where the R⁷ or R⁹ groups are NH₂ to afford analogs of (XXVIII) where the alkylamino group is on the R⁷ or R⁹ position.

SCHEME 10

An alternate method for preparing secondary anilines (XXVI) or α-substituted secondary anilines (XXVIII) proceeds from bromides (XIX). Treatment of bromide (XIX) with a variety of alkyl or benzylamines (XXIX), which can be chiral if R^{12a} and R^{12a} are appropriate groups, in the presence of a Pd(0) catalyst, such as Pd₂(dba)₃, Pd(PPh₃)₄ or Pd(PPh₃)₂Cl₂, and suitable ligand such BINAP or PPh₃, and a base such as NaO_tBu or CsCO₃ in a suitable solvent such as DMF, toluene, THF, DME, or the like, affords the anilines (XXVIII). In analogy of Scheme 8, the protocol described in Scheme 11 can also be applied to analogs of (XIX) where the R⁷ or R⁹ groups are Br, I, OTf, etc., to afford analogs of (XXX) or where the alkylamino group is on the R⁷ or R⁹ position.

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SCHEME 11

Scheme 12 shows another example for functionalizing arylbromides. Treatment of the arylbromide (XIX) with suitable base such as n-BuLi or t-BuLi followed by addition of B(O-*i*Pr)3 in a suitable solvent such as THF, DME, or the like, affords an aryl boronic ester intermediate. Treatment of the intermediate with suitable acid such as HOAc followed by oxidation with H₂O₂ affords the phenol derivatives (XXX). Coupling of these phenols (XXX) with various alkylhalides or alkylsulfonates (XXXI) in the presence of a suitable base such as NaH or KOH in a suitable solvent such as DMF, DMSO, toluene, THF, DME, or the like, affords the alkoxy indoline (XXXII). In analogy of Scheme 8, the protocol described in Scheme 12 can also be applied to analogs of (XIX) where the R⁷ or R⁹ groups are Br, I, OTf, etc., to afford analogs of (XXXII) where the alkoxy group is on the R⁷ or R⁹ position.

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SCHEME 12

Br
$$R^9$$
 R^{1} $R^{$

Alternatively, various alcohols (XXXIII) couple to the phenols (XXX) under Mitsunobu reaction condition (See Mitsunobu, O. *Synthesis* **1981**, 1-28) in the presence of diethylazodicarboxylate (DEAD) with a suitable ligand such as PPh3 or Et3P in a suitable solvent such as THF to afford the alkoxy indoline (XXXII) as shown in Scheme 13. In analogy of Scheme 8, the protocol described in Scheme 13 can also be applied to analogs of (XXX) where the R⁷ or R⁹ group is OH to afford analogs of (XXXII) or where the alkoxy group is on the R⁷ or R⁹ position.

SCHEME 13

In addition, the phenols (XXX) also reacts with a functionalized aryl boronic acid (XXXIV) in the presence of Cu(II) species, such as Cu(OAc)₂ or CuF₆(MeCN)₄ and a base such as NEt₃ or K₂CO₃ in a suitable solvent such as CH₂Cl₂ to afford the

aryloxy indoline (XXXV) as shown in Scheme 14. In analogy of Scheme 8, the protocol described in Scheme 14 can also be applied to analogs of (XXX) where the R^7 or R^9 group is OH to afford analogs of (XXXV) or where the aryloxy group is on the R^7 or R^9 position.

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SCHEME 14

HO
$$\mathbb{R}^9$$
 \mathbb{R}^7 \mathbb{R}^1 \mathbb{R}^4 \mathbb{R}^4

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Similarly arylbromides can be converted to thiophenol derivatives as shown in Scheme 15. Treatment of the arylbromide (XIX) with suitable base such as n-BuLi or t-BuLi followed by addition of sulfur in a suitable solvent such as pentane, hexane, THF, DME, or the like, followed by aqueous work-up affords the thiophenol (XXXVI). Various alkylhalides or alkylsulfonates (XXXI)can be coupled to the thiophenol (XXXVI) in the presence of a suitable base such as K2CO3, Na2CO3, NaH or KOH in a suitable solvent such as DMF, DMSO, toluene, THF, DME, or the like, affords the sulfide derivatives (XXXVII). These sulfides (XXXV) can be oxidized by suitable oxidizers such as MCPBA, NaIO4, H2O2, KMnO4, or oxone in a suitable solvent such as CH2Cl2, CHCl3, MeOH, EtOH, H2O, or the like to give sulfoxides or sulfones (XXXVIII). In analogy of Scheme 8, the protocol described in Scheme 15 can also be applied to analogs of (XIX) where the R⁷ or R⁹ groups are Br, I, OTf, etc., to afford analogs of (XXXVII) and (XXXVIII) where the alkylthio, alkylsulfoxy or alkylsulfonyl group is on the R⁷ or R⁹ position.

SCHEME 15

Br
$$R^9$$
 R^{1} $R^{$

Another example is shown in Scheme 16. Treating bromide derivatives (XIX) with an appropriate alkyl zinc reagent (XXXIX), which can be generated from the corresponding alkyl halide, in the presence of Pd(0) catalyst such as Pd2(dba)3, Pd(PPh3)4 or Pd(PPh3)2Cl2, and with or without a copper(I) salt, affords the derivatives (XL) where R⁸ is a alkyl group (see Knochel, P., et. al. *Chem. Rev.* 1993, 93, 2117; and Weichert, A., et. al. *Syn. Lett.* 1996, 473). In analogy of Scheme 8, the protocol described in Scheme 16 can also be applied to analogs of (XIX) where the R⁷ or R⁹ groups are Br, I, OTf, etc., to afford analogs of (XL) where the alkoxy group is on the R⁷ or R⁹ position.

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SCHEME 16

Furthermore, as an extension of this approach to a rapid preparation of a large array of biaryl indole and indoline derivatives, these bromide derivatives (XIX or XXIV) can be bound to a solid support and the arylamine couplings can be carried out on solid support as illustrated in Scheme 17. Towards that end treatment of indoline (XIX) with TFA in CH2Cl2, to remove the Boc protecting group, followed by extraction from aqueous base provides the free amine (XIX). The free amine can be loaded onto a suitable solid support such as (XLI) using conditions well known to those skilled in the art. Thus, p-nitrophenylchloroformate Wang resin (XLI) which can be obtained commercially from sources such as Novabiochem, Inc. is swollen in a suitable solvent such as N-methyl pyrrolidinone and treated with 1.5 equiv. of amine to afford the functionalized resin (XLII). Arylamine couplings are then carried out in array format by treatment of resins (XLII) with a suitable palladium source such as Pd2(dba)3, Pd(PPh3)4, Pd(dppf)Cl2 or Pd(PPh3)2Cl2, and suitable ligand such BINAP or PPh3, and a base such as NaOtBu or CsCO3 with an excess (typically 5 equivalents) of an aniline. The coupling may be repeated to ensure complete conversion to the desired coupled product. Cleavage from the solid support by treatment with TFA affords the corresponding indoles and indolines (XLIII) as their TFA salts.

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SCHEME 17

In addition, derivatives of type (I) can be alkylated with any number of functionalized alkyl sidechains. Typical procedures utilizing standard alkylation of a secondary amine with an alkylhalide under base catalyzed conditions are well known by those skilled in the art. For example, the secondary amino group of Formula (I) $(R^1 = H)$ can be alkylated with alkylhalides or alkylsulfonates in the presence of NaI or KI and a base such as K_2CO_3 , Na_2CO_3 , triethylamine, or the like, in dioxane or THF or other such solvent while heating (see Glennon, R.A., et. al., *Med. Chem. Res.*, 1996, 197) affords the R^1 alkylated indolines.

It is understood that the compounds of the present invention can be prepared in a number of ways well known to one skilled in the art of organic synthesis. The compounds of the present invention can be synthesized using the methods described herein, together with synthetic methods known in the art of synthetic organic chemistry, or variations thereon as appreciated by those skilled in the art. Additional methods include, but are not limited to, those described in USSN 09/594,954 (filed

June 15,2000); USSN 09/595,250 (filed June 15, 2000); USSN 09/594,008 (filed June 15, 2000); USSN 10/026,793 (filed December 19, 2001); USSN 10/026,611 (filed December 19, 2001); USSN 10/026,404 (filed December 19, 2001); wherein all references are hereby incorporated in their entirety herein by reference.

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EXAMPLES

EXAMPLE 1

5 cis-(4a,9b)-8-bromo-6-methylsulfanyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole

Step A. 2-Methylsulfanyl-phenylamine (23.6 g, 169.5 mmol) was suspended in conc. HCl (200 mL) and trifluoroacetic acid (130 mL), and cooled to 0 °C in an ice bath. Sodium nitrite (14.0 g, 203.4 mmol) was dissolved in water (45 mL) and added dropwise to the suspension over 45 min. After the addition, the reaction was stirred at 0 °C for 1 h. In a separate flask, tin(II) chloride (76 g, 338.4 mmol) was dissolved in conc. HCl (100 mL) and added slowly over 15 min to the reaction mixture. The resultant suspension was warmed to rt and stirred for 48 h. The reaction was filtered, washed with *iso*-propyl alcohol (15 mL), and dried to give 1-(2-methylsulfanyl-phenyl)hydrazine hydrochloride (30 g, 93%). ¹H NMR (CD3OD, 300 MHz) δ 7.44 (d, 1H, J = 7.7 Hz), 7.32-7.26 (m, 1H), 7.08-7.00 (m, 2H), 2.39 (s, 3H) ppm.

Step B. 1-(2-Methylsulfanyl-phenyl)hydrazine hydrochloride (27 g, 141.9 mmol) and 4-piperidone monohydrate hydrochloride (21.8 g, 141.9 mmol) were dissolved in ethanol (350 mL) and heated at reflux for 45 min. Conc. HCl (30 mL) was added and heated at reflux for 12 h. The reaction was cooled to rt, filtered, washed with cold isopropyl alcohol (50 mL), and dried to give an off white solid, 6-methylsulfanyl-2,3,4,5-tetrahydro-1*H*-pyrido[4,3-*b*]indole hydrochloride (33 g, 92%). 1 H NMR (CD₃OD, 300 MHz) δ 7.24 (d, 1H, J = 8.7 Hz), 7.08 (dd, 1H, J = 0.7, 7.3 Hz), 6.93 (t, 1H, J = 7.7 Hz), 3.29-3.27 (m, 2H), 3.17 (t, 2H, J = 5.85 Hz), 2.85 (t, 2H, J = 5.65 Hz), 2.44 (s, 3H) ppm.

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Step C. 6-Methylsulfanyl-2,3,4,5-tetrahydro-1*H*-pyrido[4,3-*b*]indole hydrochloride (20 g, 78.74 mmol) was suspended in trifluoroacetic acid (562 mL) and cooled to 0 °C in an ice bath. NaCNBH3 (19.53 g, 314.96 mmol) was added portion wise over 25 min and the mixture was stirred at 0 °C for 4 h. The reaction mixture was basified to pH 10 with conc. ammonium hydroxide and extracted with ethyl acetate (4 x 500 mL) and the organic layer separated. The organics were collected, dried over magnesium sulfate, and filtered to give *cis*-(4a,9b)-6-methylsulfanyl-2,3,4,4a,5,9b-hexahydro-1*H*-pyrido[4,3-*b*]indole. ¹H NMR (CDCl3, 300 MHz) δ 7.11 (d, 1H, J = 7.7 Hz), 6.99 (d, 1H, J = 7.3 Hz), 6.73 (t, 1H, J = 5.7 Hz), 4.11 (s, 1H), 3.95-3.92 (m, 2H), 3.27-3.01 (m, 3H), 2.93-2.83 (m, 2H), 2.41 (s, 3H), 2.02-1.91 (m, 1H), 1.82-1.74 (m, 1H) ppm.

Step D. *Cis*-(4a,9b)-6-methylsulfanyl-2,3,4,4a,5,9b-hexahydro-1*H*-pyrido[4,3-*b*]indole (4 g, 18.2 mmol) was dissolved in CH₂Cl₂ (30 mL) and saturated potassium carbonate (25 mL)was added. This mixture was stirred vigorously and cooled to 0 °C in an ice bath. Di-*tert*-butyl dicarbonate (3.9 g, 18.2 mmol) and 4-dimethylamino-pyridine (10 mg, 0.082mmol) in CH₂Cl₂ (20 mL) were added in 5 portions over 15 min. The mixture was stirred at 0 °C for 2 h. The organic layer was separated and aqueous layer was back extracted with CH₂Cl₂ (2 x 150 mL). The combined organics were dried over magnesium sulfate, filtered and concentrated under reduced pressure to give *tert*-butyl *cis*-(4a,9b)-6-methylsulfanyl-1,3,4,4a,5,9b-hexahydro-2*H*-pyrido[4,3-*b*]indole-2-carboxylate (4.84 g, 83%). ¹H NMR (DMSO d₆ 300 MHz) δ 6.99 (d, 1H, J = 7.8 Hz), 6.93 (dt, 1H, J = 2.2, 7.4 Hz), 6.56 (t, 1H, J = 7.6 Hz), 4.00-3.95 (m, 1H), 3.59 (dd, 1H, J = 5.3, 8.3 Hz), 3.45-3.26 (m, 4H), 2.34 (s, 3H), 1.90-1.84 (m, 1H), 1.71-1.69 (m, 1H), 1.36 (s, 9H) ppm.

Step E. Tert-butyl cis-(4a,9b)-6-methylsulfanyl-1,3,4,4a,5,9b-hexahydro-2H-pyrido[4,3-b]indole-2-carboxylate (1 g, 3.13mmol) was dissolved in DMF (4 mL) at rt, and cooled to 0 °C in an ice bath. In a separate flask N-bromosuccinimide (NBS) (0.56 g, 3.13 mmol) was dissolved in DMF (1.5 mL) and added slowly to the first solution over 5 min. The reaction mixture was stirred for 1 h at 0 °C. The reaction was followed by thin layer chromatography until no more starting material was present. The reaction mixture was quenched with water (30 mL) and extracted with ethyl acetate (3 x 40 mL). The organic layers were collected and washed with brine

(3 x 34 mL) and water (1 x 30 mL). The organic layer was dried over magnesium sulfate, filtered, and concentrated under reduced pressure to give an oil. The oil was purified by silica gel column chromatography, (15% ethyl acetate, hexanes) to afford *tert*-butyl *cis*-(4a,9b)-8-bromo-6-methylsulfanyl-1,3,4,4a,5,9b-hexahydro-2*H*-pyrido[4,3-*b*]indole-2-carboxylate as an oil (0.64 g, 51%). ¹H NMR (CDCl₃, 300 MHz) δ 7.18 (d, 1H, J = 1.9 Hz), 7.11 (s, 1H), 4.09-4.00 (m, 1H), 3.74-3.51 (m, 3H), 3.39-3.28 (m, 2H), 2.41 (s, 3H), 1.98-1.84 (m, 1H), 1.75-1.60 (m, 1H), 1.47 (s, 9H) ppm.

Step F. *Tert*-butyl *cis*-(4a,9b)-8-bromo-6-methylsulfanyl-1,3,4,4a,5,9b-hexahydro-2*H*-pyrido[4,3-*b*]indole-2-carboxylate (25 mg, 0.063 mmol) was dissolved in trifluoroacetic acid/chloroform (1:2, 1.5 mL) and stirred for 0.5 h. The solvent was evaporated under reduced pressure, and the residue was dissolved in water. The aqueous solution was basified with concentrated ammonium hydroxide to pH 12, and extracted with chloroform (3 x 10 mL). The organic solution was washed with brine, dried over magnesium sulfate, filtered and concentrated under reduced pressure to give the title compound as a yellow semi-solid (17.7 mg, 95%). ¹H NMR (300MHz, CD₃OD) δ 7.27 (d, 1H, J = 2.2 Hz), 7.18 (d, 1H, J = 2.2 Hz), 3.50-3.10 (m, 3H), 3.11 (s, 3H), 2.68 (dd, 1H, J = 12.9, 9.6 Hz), 2.38, (s, 3H), 2.28-2.05 (m, 2H) ppm MS (EI) 315.0 (M+H).

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EXAMPLE 2

cis-(4a,9b)-8-bromo-5-methyl-6-(methylsulfanyl)-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole

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Step A. To a methylene chloride (300 mL) solution of 6-methylsulfanyl-2,3,4,5-tetrahydro-1*H*-pyrido[4,3-*b*]indole hydrochloride (from Example 1, Step B)

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(18.03 g, 71 mmol) was added triethylamine (14.3 g, 142 mmol) in one portion and resulted solution was stirred at rt for 10 min. The reaction was cooled to 0° C in an ice bath then di-*tert*-butyl dicarbonate (18.6 g, 85 mmol) was added, and then reaction was allowed to warm slowly to rt and stir for 14 h. Reaction mixture was poured into water (300 mL) and then layers separated. The aqueous layer was extracted with chloroform (3 x 100 mL) and the organics collected, washed with brine (150 mL), dried over MgSO4, and concentrated to dryness under reduced pressure. The resultant semi-solid was recrystalized from ethyl acetate to give *tert*-butyl 6-methylsulfanyl-1,3,4,5-tetrahydro-2*H*-pyrido[4,3-*b*]indole-2-carboxylate as sandy brown solid (7.2 g, 32%). ¹H NMR (CDCl₃, 300 MHz) δ 8.19 (br s, 1H), 7.36 (d, 1H, J = 8.0 Hz), 7.25 (d, 1H, J = 6.9 Hz), 7.09 (t, 1H, J = 7.7 Hz), 4.63 (br s, 2H), 3.82 (br s, 2H), 2.91-2.82 (m, 2H), 2.50 (s, 3H), 1.50 (s, 9H) ppm.

Step B. *Tert*-butyl 6-methylsulfanyl-1,3,4,5-tetrahydro-2*H*-pyrido[4,3-*b*]indole-2-carboxylate (7.2 g, 22.6 mmol), potassium hydroxide (6.3g, 113 mmol), and iodomethane (32 g, 226 mmol) were combined with dry DME (110 mL) and stirred at rt for 10 h. The reaction was then filtered and the residue washed with chloroform. The filtrate was concentrated under reduced pressure to give *tert*-butyl 5-methyl-6-methylsulfanyl-1,3,4,5-tetrahydro-2*H*-pyrido[4,3-*b*]indole-2-carboxylate as an oily brown residue (7.7 g, 105%). ¹H NMR (CDCl₃, 300 MHz) δ 7.32 (d, 1H, J = 7.7 Hz), 7.12 (d, 1H, J = 6.6 Hz), 7.03 (t, 1H, J = 7.0 Hz), 4.61 (s, 2H), 4.08 (s, 3H), 3.85 (br s, 2H), 2.80 (br s, 2H), 2.49 (s, 3H), 1.49 (s, 9H) ppm

Step C. Tert-butyl 5-methyl-6-methylsulfanyl-1,3,4,5-tetrahydro-2H-pyrido[4,3-b]indole-2-carboxylate (7.7 g, 23.9 mmol) was stirred in trifluoroacetic acid (120 mL) at 0°C for 10 min, then sodium cyanoborohydride (8.25 g, 119 mmol) was added slowly while monitoring the internal reaction temperature to be below 5°C during the addition. After the addition, the reaction was allowed to warm to rt under nitrogen and stir for 2 h. Ice chips were added and the reaction was cooled in an ice bath, and then basified to a pH~13 with 50% sodium hydroxide. The aqueous mixture was then extracted with chloroform (3 x 80 mL) and then the organics were collected, washed with brine (1x 100 ml), dried over magnesium sulfate, and concentrated under reduced pressure to an oily residue. The residue was dissolved in chloroform (75 mL) and to this was added di-tert-butyl dicarbonate (3.2 g) and triethylamine (3.0 mL) the

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reaction was stirred for 2 h. The solvent evaporated under reduced pressure to give an oil. This oil was purified by silica gel column chromatography eluting with (5%, then 10%) ethyl acetate/hexanes, to give *tert*-butyl *cis*-(4a,9b)-5-methyl-6-methylsulfanyl-1,3,4,4a,5,9b-hexahydro-2*H*-pyrido[4,3-*b*]indole-2-carboxylate as a colorless oil (2.85 g, 36%) ¹H NMR (CDCl₃, 300 MHz) δ 7.13 (dd, 1H, J = 7.7, 1.1 Hz), 6.96 (d, 1H, J = 7.4 Hz), 6.70 (t, 1H, J = 7.3 Hz), 4.00-3.27 (m, 6H), 3.14 (s, 3H), 2.36 (s, 3H), 1.98-1.85 (m, 2H0, 1.42 (s, 9H) ppm.

Step D. *Tert*-butyl *cis*-(4a,9b)-5-methyl-6-methylsulfanyl-1,3,4,4a,5,9b-hexahydro-2*H*-pyrido[4,3-*b*]indole-2-carboxylate (60 mg, 0.19 mmol) was dissolved in anhydrous DMF (1 mL) at 0°C, to this was added NBS (33 mg, 0.19 mmol) in DMF (0.5 mL). This was stirred at 0°C for 1 h. Then water (3 mL) was added and reaction partitioned over chloroform. Layers were separated and aqueous layer extracted with chloroform (3x 5 mL). The organics were collected, washed with water (3x 10 mL), dried over magnesium sulfate, and concentrated under reduced pressure to *tert*-butyl *cis*-(4a,9b)-8-bromo-5-methyl-6-methylsulfanyl-1,3,4,4a,5,9b-hexahydro-2*H*-pyrido[4,3-*b*]indole-2-carboxylate as a yellow solid (57 mg, 76%). ¹H NMR (CDCl₃, 300 MHz) δ 7.19 (d, 1H, J = 1.8 Hz), 7.02 (d, 1H, J = 1.9 Hz), 3.60-3.20 (m, 6H), 3.10 (s, 3H), 2.36 (s, 3H), 1.94-1.80 (m, 2H), 1.43 (s, 9H) ppm.

Step E. *Tert*-butyl *cis*-(4a,9b)-8-bromo-5-methyl-6-methylsulfanyl-1,3,4,4a,5,9b-hexahydro-2*H*-pyrido[4,3-*b*]indole-2-carboxylate (20 mg, 0.048 mmol) was dissolved in trifluoroacetic acid/chloroform (1:2, 1.5 mL) and stirred for 0.5 h. The solvent was evaporated under reduced pressure to give the title compound as a yellow semi-solid (19 mg, 90%). ¹H NMR (CD₃OD, 300 MHz) δ 7.27 (d, 1H, J = 2.2 Hz), 7.17 (d, 1H, J = 2.2 Hz), 3.50-3.30 (m, 3H), 3.28-3.12 (m, 2H), 3.11 (s, 3H), 2.69 (dd, 1H, J = 9.6, 3.3 Hz), 2.38 (s, 3H), 2.28-2.00 (m, 2H) ppm. MS (CI, NH3) 315 (M + H).

EXAMPLE 3

cis-(4a,9b)-6-methylsulfanyl-2,3,4,4a,5,9b-hexahydro-1*H*-pyrido[4,3-*b*]indol-8-amine

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Step A. *Tert*-butyl *cis*-(4a,9b)-8-bromo-6-methylsulfanyl-1,3,4,4a,5,9b-2*H*-pyrido[4,3-*b*]indole-2-carboxylate (from Example 1 step E)(950 mg, 2.37 mmol) and di-*tert*-butyl dicarbonate (1.29 g, 5.92 mmol) were mixed together neat and heated at 86 °C for 6.5 h under nitrogen. The resultant residue was purified by silica gel column eluting with 7.5% ethylacetate/hexanes to give di(*tert*-butyl) *cis*-(4a,9b)-8-bromo-6-methylsulfanyl-3,4,4a,9b-tetrahydro-1*H*-pyrido[4,3-*b*]indole-2,5-dicarboxylate as a foam (1.13 g, 95%). ¹H NMR (DMSO d₆, 300 MHz) δ 7.29 (s, 1H), 7.19 (s, 1H), 4.69-4.61 (m, 1H), 4.22-4.18 (m,1H), 3.71-3.62 (m,1H), 3.57-3.37 (m, 2H), 2.98-2.84 (m, 2H), 2.37 (s, 3H), 2.00-1.96 (m, 1H), 1.51 (s, 9H), 1.37 (s, 9H) ppm.

Step B. Di(tert-butyl) cis-(4a,9b)-8-bromo-6-methylsulfanyl-3,4,4a,9btetrahydro-1*H*-pyrido[4,3-*b*]indole-2,5-dicarboxylate (100 mg, 0.20 mmol), 20 benzophenone imine (43 mg, 0.24 mmol), and sodium tert-butoxide (29 mg, 0.30 mmol) were dissolved/suspended in anhydrous toluene (1.5 mL) and then argon was bubbled through the reaction solution for 10 min. Then (R)-(+)-2,2'bis(diphenylphosphino)-1,1'-binaphthyl (5 mg, 0.008 mmol) and tris(dibenzylideneacetone)dipalladium (0) (2 mg, 0.002 mmol) were added and then argon was bubbled through the reaction solution for 10 min, then heated at 90 °C for 5 25 h. Reaction mixture was cooled to rt, diluted with ethyl acetate (25 ml) and filtered. The filtrate was concentrated under reduced pressure to give a yellow colored oil. The oil was dissolved in methyl alcohol and to it added sodium acetate (33 mg, 0.40 mmol) and hydroxylamine hydrochloride (42 mg, 0.60 mmol) and stirred for 20 min 30 at rt. Reaction was quenched with 6N sodium hydroxide to pH ~ 12, and extracted

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with ethyl acetate (3 x 15 mL). Organic layer was separated, dried over magnesium sulfate, and concentrated under reduced. The oil residue was purified by silica gel column chromatography to give di(*tert*-butyl) *cis*-(4a,9b)-8-amino-6-methylsulfanyl-3,4,4a,9b-tetrahydro-1*H*-pyrido[4,3-*b*]indole-2,5-dicarboxylate as a foam (37 mg, 42%). 1 H NMR (CDCl₃, 300 MHz) δ 6.46-6.36 (m, 2H), 4.69-4.59 (m, 1H), 4.38-4.24 (m, 1H), 3.91-3.82 (m, 1H), 3.72-3.20 (m, 4H), 3.00-2.78 (m, 2H), 2.40 (s, 3H), 2.05-1.92 (m, 1H), 1.60-1.34 (m, 18 H) ppm.

Step C. Di(*tert*-butyl) *cis*(4a,9b)-8-amino-6-methylsulfanyl-3,4,4a,9b-tetrahydro-1*H*-pyrido[4,3-*b*]indole-2,5-dicarboxylate (37 mg, 0.085 mmol) was dissolved in chloroform (3 mL) and ethyl alcohol (0.4 mL) at rt. A stream of dry hydrogen chloride gas was allowed to bubble through this solution for 3 min. The reaction was concentrated under reduced pressure to give the title compound as a white solid (26 mg, 90%). ¹H NMR (CD₃OD, 300 MHz) δ 7.14 (s, 2H), 4.07-4.02 (m, 1H), 3.58-3.40 (m, 2H), 2.89-2.85 (m, 1H), 2.46 (s, 3H), 2.19-2.14 (m, 2H) ppm. MS (ESI): 236.3 (M+H).

EXAMPLE 4

[cis-(4a,9b)-6-methylsulfanyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl]-phenyl-amine

Step A. Triphenylbismuth (0.220 g, 0.50 mmol) was stirred with iodobenzene diacetate (0.18 g, 0.55 mmol) in CH₂Cl₂ (5 mL) for 15 h at rt. The solvent was evaporated and Et₂O (2 mL) with heptane (2 mL) was added and heated. The resulting solid was filtered hot affording bis(acetato)trisphenylbismuth (0.19 g, 70%) as a white flaky solid. ¹H NMR (CDCl₃, 300 MHz) δ 8.15 (dd, 6H, J = 1.1, 8.4 Hz), 7.62 (t, 6H, J = 7.3 Hz), 7.50 (t, 3H, J = 8.4 Hz), 1.81 (s, 6H) ppm.

- **Step B.** Di(*tert*-butyl) *cis*-(4a,9b)-8-amino-6-methylsulfanyl-3,4,4a,9b-tetrahydro-1*H*-pyrido[4,3-*b*]indole-2,5-dicarboxylate (from example 6 step A 76 mg, 0.152 mmol) was combined with bis(acetato)trisphenylbismuth (93 mg, 0.167 mmol) and copper(II) acetate (2 mg, 0.015 mmol) in CH₂Cl₂ (2 mL) and stirred for 1.5 h.
- The solvent was evaporated and the black colored residue purified by silica gel column chromatography (20% EtOAc/Hex), affording a clear oil. This oil was then dissolved in EtOH (0.5mL) and chloroform (2 ml) and HCl gas was allowed to bubble through for 10 min. The solvent was evaporated to give the title compound (50 mg, 78%). ¹H NMR (CD₃OD, 300 MHz) δ 7.37-7.32 (m, 2H), 7.29-7.18 (m, 2H), 7.166.97 (m, 3H), 4.51-4.42 (m, 1H), 3.82-3.58 (m, 2H), 3.40-3.24 (m, 3H), 2.54 (s, 3H), 2.48-2.10 (m, 2H) ppm.

EXAMPLE 5

15 (4-fluorophenyl)-[cis-(4a,9b)6-methylsulfanyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl]-amine

Step A. To a solution of *p*-fluorophenylmagnesium bromide (48.8 mL, 1.0 M THF) in dry diethyl ether (20mL) was added bismuth(III)chloride (5.0 g, 15.8 mmol). The reaction was allowed to proceed at rt for 2 h. Ice was added and the aqueous layer extracted with ether (3 x 25 mL). The combined extracts were washed with brine (25 mL) and dried (MgSO4) and evaporated. The residue was combined iodobenzene diacetate (3.4 g, 10.5 mmol) in CH₂Cl₂ (20 mL) for 15 h at rt. The solvent was evaporated affording bis(acetato)tris(4-fluorophenyl)bismuth (3.28 g, 34%). ¹H NMR (CDCl₃, 300 MHz) δ 8.15–8.21 (m, 6H), 7.20–7.28 (m, 6H), 1.81 (s, 6H) ppm.

Step B. The title compound was prepared according to the procedure of Example 25, Step B (49 mg, 73%) using bis(acetato)tris(4-fluorophenyl)bismuth. ¹H NMR (CD₃OD, 300 MHz) δ 7.20-6.93 (m, 6H), 4.53-4.40 (m, 1H), 3.82-3.56 (m, 2H), 3.40-3.31 (m, 3H), 2.53 (s, 3H), 2.50-2.01 (m, 2H) ppm.

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EXAMPLE 6

(4-methoxy-2-methyl-phenyl)-[cis-(4a,9b)-6-methylsulfanyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl]-amine

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Step A. To a solution of 1-bromo-4-methoxy-2-methylbenzene (0.58 mg, 3.0 mmol) in dry THF (15mL) was added magnesium turnings. The reaction was heated at 60 °C in an oil bath and the magnesium turnings were gently crushed with a glass rod. A small crystal of iodine was also added. Reaction was removed from the oil bath when bubbling was observed, and 1-bromo-4-methoxy-2-methylbenzene (5.18 g, 27.00 mmol) was added slowly in dry THF (25 mL). Reaction was stirred at rt for 15 min. Then the solution was transferred via syringe to a dry flask under nitrogen and cooled to 0 °C in an ice bath. Bismuth(III)chloride (2.84 g, 315 mmol) was added slowly and the resultant heterogeneous solution was allowed to warm to rt and stir for 4 h. The reaction was filtered through a bed of Celite and the filtrate collected. The filtrate was poured onto ice and the aqueous layer extracted with ethyl acetate (2 x 50 mL). The combined extracts were washed with brine (25 mL) and dried (MgSO₄) and evaporated. The residue was recrystallized from ethyl acetate to give a yellowish solid. This solid was combined with iodobenzene diacetate (1.33 g, 4.1 mmol) in CH2Cl2 (45 mL) for 15 h at rt. The solvent was evaporated affording bis(acetato)tris(2-methyl-4-methoxyphenyl)bismuth (805 mg, 4.3 %). ¹H NMR

(CDCl₃, 300 MHz) δ 8.21(d, 3H, J = 8.5 Hz), 6.95 (s, 6H), 3.83 (s, 9H), 2.57 (s, 9H), 1.73 (s, 6H) ppm.

Step B. The title compound was prepared according to the procedure of Example 7, Step B (49 mg, 73%) using bis(acetato)tris(2-methyl-4-methoxyphenyl)bismuth. ¹H NMR (CD₃OD, 300 MHz): δ 7.10-7.05 (m, 1H), 6.84 (s, 1H), 6.80-6.72 (m, 1H), 6.64 (s, 1H), 6.51 (s, 1H), 4.52-4.41 (m, 1H), 3.76 (s, 3H),

3.66-3.56 (m, 1H), 3.40-3.19 (m, 3H), 2.48-2.39 (m, 4H), 2.17-2.04 (m, 4H) ppm.

EXAMPLE 7

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trans-(4a,9b)-8-bromo-2,5-dimethyl-6-[(4-methylphenyl)sulfanyl]-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole

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Step A. 2-Iodoaniline (16 g, 73 mmol) was suspended in concentrated hydrochloric acid (100 mL), and then cooled to 0 °C in an ice bath. Sodium nitrite (6 g, 87.6 mmol) in water (25 mL) was added slowly to reaction mixture and then reaction allowed to stir at 0 °C for 1.5 h. In a separate flask, tin(II) chloride (84.7 g, 182.5 mmol) was dissolved in concentrated hydrochloric acid (12 mL) and added slowly over 30 min to reaction mixture. The resulting suspension was allowed to warm to rt and stirred for 14 h. The solid was filtered off, and allowed to dry to afford 1-(2-iodophenyl)hydrazine hydrochloride (19 g, 96%). ¹H NMR (CDCl₃, 300 MHz) δ 7.82 (dd, 1H, J = 1.1, 7.7 Hz), 7.39 (dt, 1H, 1.2, 7.7 Hz), 6.96 (dd, 1H, J = 1.1, 8.1 Hz), 6.82 (dt, 1H, J = 1.1, 7.5) ppm. MS (ApCl) 275 (M⁺+CH3CN+H).

Step B. 1-(2-Iodophenyl)hydrazine hydrochloride (1.67 g, 6.2 mmol) and 4-piperidone monohydrate hydrochloride (0.952 g, 6.2 mmol) were dissolved in

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trifluoroethanol (15 mL) and concentrated hydrochloric acid (5 mL) and heated at 87 °C and stirred for 3 h. The solid was filtered, washed with cold isopropyl alcohol (50 mL), and dried to give 6-iodo-2,3,4,5-tetrahydro-1H-pyrido[4,3-b]indole hydrochloride (1.76 g, 85%) as a tan solid. ¹H NMR (CD₃OD, 300 MHz) δ 7.49 (d, 1H, J = 7.7 Hz), 7.43 (d, 1H, J = 8 Hz), 6.82 (t, 1H, J = 7.7 Hz), 4.40 (s, 2H), 3.60 (t, 2H, J = 6.25 Hz), 3.18 (t, 2H, J = 5.85 Hz) ppm.

Step C. 6-Iodo-2,3,4,5-tetrahydro-1H-pyrido[4,3-b]indole hydrochloride (7g, 21 mmol) was suspended in CH₂Cl₂ (150 mL) and to this suspension was added 4-dimethylamino-pyridine (0.1 g, 0.82 mmol) and saturated potassium carbonate solution (150 mL) with stirring. Then di-tert-butyl dicarbonate (5.5 g, 25.2 mmol) in dichloromethane (20 mL) was added in 5 portions over 5 min. The resulting two-phase mixture was stirred vigorously at rt for 1.5 h. Layers were separated and aqueous layer was back extracted with CH₂Cl₂ (2 x 100 mL). The organics were collected, dried over magnesium sulfate, filtered and then concentrated under reduced pressure to give tert-butyl 6-iodo-1,3,4,5-tetrahydro-2H-pyrido[4,3-b]indole-2-carboxylate (7.06 g, 84%). ¹H NMR (CDCl₃ 300 MHz) δ 7.63 (d, 1H, J = 7.7 Hz), 7.38 (d, 1H, J = 7.7 Hz), 6.76 (t, 1H, J = 7.5 Hz), 4.59 (br s, 2H), 3.82 (br s, 2H), 2.78 (t, 2H, J = 5.3 Hz), 1.49 (s, 9H) ppm.

Step D. Tert-butyl 6-iodo-1,3,4,5-tetrahydro-2H-pyrido[4,3-b]indole-2-carboxylate (6.8 g, 17 mmol) was dissolved in DME (50 mL), and potassium hydroxide (4.8 g, 85.4 mmol) and iodomethane (15.7 g, 110.5 mmol) were added and heated at 80 °C in a pressure vessel for 3 h. The reaction was cooled to rt and diluted with ethyl acetate (50 mL). The solids were removed by vacuum filtration. The filtrate was concentrated under reduced pressure to give a brown oil (5.5 g, 79% crude yield). The oil was dissolved in CH₂Cl₂ (30 mL). Trifluoroacetic acid (30 mL) was added in ten portions over 5 min and stirred for 30 min. The reaction was basified with 50% sodium hydroxide to pH 12. This mixture was extracted with CH₂Cl₂ (3 x 150 mL). The organics were collected, dried over magnesium sulfate, filtered and then concentrated under reduced pressure to give 6-iodo-5-methyl-2,3,4,5-tetrahydro-1H-pyrido[4,3-b]indole (3.85 g, 73%). ¹H NMR (CDCl₃ 300 MHz) δ 7.62 (d, 1H, J

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= 7.6 Hz), 7.33 (d, 1H, J = 7.7 Hz), 6.74 (t, 1H, J = 7.5 Hz), 4.03 (s, 2H), 3.95 (s, 3H), 3.26 (t, 2H, J = 5.6 Hz), 2.71 (t, 2H, J = 5.6 Hz) ppm.

Step E. 6-Iodo-5-methyl-2,3,4,5-tetrahydro-1H-pyrido[4,3-b]indole (3.85 g, 12.3 mmol) was suspended in methanol (40 mL). Formaldehyde (14 mL of 37%) was added and heated at reflux for 2 h. The reaction was cooled to 0 °C in an ice bath and sodium borohydride (1.7 g, 46 mmol) was added slowly over 15 min and stirred for 2 h at 0-10 °C. The reaction was diluted with water (200 mL) and extracted with CH₂Cl₂ (3 x 150 mL). The organics were collected, washed with brine (250 mL), dried over magnesium sulfate, filtered and concentrated under reduced pressure. The product was purified by silica gel column chromatography (3% MeOH/CH₂Cl₂) to give 6-iodo-2,5-dimethyl-2,3,4,5-tetrahydro-1H-pyrido[4,3-b]indole (0.6 g, 15%). 1 H NMR (CDCl₃ 300 MHz) δ 7.59 (d, 1H, J = 7.4 Hz), 7.33 (d, 1H, J = 7.6 Hz), 6.73 (t, 1H, J = 7.7 Hz), 3.98 (s, 3H), 3.62 (s, 2H), 2.83 (s, 4H), 2.55 (s, 3H) ppm.

Step F. 6-Iodo-2,5-dimethyl-2,3,4,5-tetrahydro-1H-pyrido[4,3-b]indole (1 g, 3 mmol) was dissolved in BH3-THF complex (15 mL) and heated at 75 °C for 18 h in a pressure vessel. The reaction was cooled to rt and concentrated under reduced pressure to a residue. The residue was heated at reflux in 6N hydrochloric acid (15 mL) for 3.5 h. The reaction was basified with 50% sodium hydroxide to pH 12. The mixture was extracted with CH2Cl2 (3 x 100 mL). The organics were collected, dried over magnesium sulfate, filtered and concentrated under reduced pressure. The residue was purified by silica gel column chromatography (4% methanol, dichloromethane) to give trans-(4a,9b)-6-iodo-2,5-dimethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole (0.12 g, 12%). ¹H NMR (CDCl3 300 MHz) δ 7.56 (dt, 1H, J = 2.2, 8.1 Hz), 7.33 (d, 1H, J = 7.3 Hz), 6.50 (t, 1H, J = 7.5 Hz), 3.48-3.41 (m, 1H), 3.04(s, 3H), 2.86-2.75 (m, 1H), 2.42 (s, 3H), 2.20-2.13 (m, 3H), 1.93-1.78 (m, 1H) ppm.

Step G. Trans-(4a,9b)-6-iodo-2,5-dimethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole (910 mg, 2.7 mmol) was combined with p-tolylthiophenol (426 mg, 3.3 mmol), sodium hydride (132 mg, 3.3 mmol, 60% in oil dispersion), copper iodide (515 mg, 2.7 mmol) and 1,3-dimethyl-3,4,5,6-tetrahydro-2(1H)-pyrimidinone (5 mL) and stirred at 100 °C for 16 h. The reaction mixture was partitioned between

water and CHCl3. The aqueous layer was extracted with CHCl3 (3 x 20 mL). The combined organics were washed with sat. NaCl (10 mL), water (10 mL), dried (MgSO4) and evaporated. The residue was loaded onto a SCX resin. The resin was washed with 150 mL of MeOH followed by washing with 2.0 M methanolic ammonia. The collected residue was dissolved in CH3CN and 1N HCl/Ether was added until no further precipitation was observed. The solid was filtered and washed with CH3CN affording the trans-(4a,9b)-2,5-dimethyl-6-[(4-methylphenyl)sulfanyl]-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole (315 mg, 36%) as a white solid. 1 H NMR (CD3OD, 300 MHz) δ 7.20 (d, 1H, J = 7.7 Hz), 7.14 (d, 1H, J = 7.3 Hz), 7.06 (d, 2H, J = 8.1 Hz), 6.97 (d, 2H, J = 8.4 Hz), 6.82–6.87 (m, 1H), 3.99–4.05 (m, 1H), 3.71–3.80 (m, 1H), 3.15–3.25 (m, 4H), 3.08 (s, 3H), 2.99 (s, 3H), 2.38–2.43 (m, 1H), 2.26 (s, 3H), 1.90–2.03 (m, 1H) ppm.

Step H. The title compound (140 mg, 45%) was prepared from trans-(4a,9b)-2,5-dimethyl-6-[(4-methylphenyl)sulfanyl]-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole (250 mg, 0.77 mmol) and NBS (164 mg, 0.92 mmol) using the same procedure described in Example 18, Step D. 1 H NMR (CD₃OD, 300 MHz) δ 7.32 (br s, 1H), 7.19 (d, 1H, J = 1.5 Hz), 7.13 (q, 4H, J= 8.4 Hz), 4.27-4.19 (m, 1H), 3.80-3.75 (m, 1H), 3.40-3.20 (m, 4H), 3.11 (s, 3H), 2.98 (s, 3H), 2.47-2.40 (m, 1H), 2.30 (s, 3H), 2.20-2.02 (m, 1H) ppm. MS (CI, NH3) 404 (M+H).

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EXAMPLE 8

cis-(4a,9b)-8-bromo-2,5-dimethyl-6-[(4-methylphenyl)sulfanyl]-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole

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Trans-(4a,9b)-8-bromo-2,5-dimethyl-6-[(4-methylphenyl)sulfanyl]-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole (Example 28, 77 mg, 0.19 mmol) and NaOMe (5 mg, 0.10 mmol) were combined with MeOH (0.5 mL) and DMF (0.5 mL). This mixture was then heated at 100 °C and MeOH was allowed to distill out.
Then CuI (3.6 mg, 0.019 mmol) was added in one portion. Reaction was maintained at 100 °C for 2 h. It was allowed to cool to rt. Ice water (10 mL) was added, solid was removed by filtration. The filtrate was extracted with chloroform (3 x 25 mL). Extracts were combined, dried, and concentrated. The residue was purified by silica gel column chromatography (CHCl3, then 3% MeOH/CHCl3) to give the title
compound (25 mg, 32%) as a semi-solid. ¹H NMR (CDCl3, 300 MHz) δ 7.26 (s, 1H), 7.11 (s, 1H), 7.07-7.04 (m, 2H), 6.99-6.96 (m, 2H), 3.33-3.19 (m, 2H), 3.02 (s, 3H), 2.67-2.60 (m, 1H), 2.44-2.37 (m, 1H), 2.29 (s, 3H), 2.26 (s, 3H), 2.23-2.19 (m, 1H), 2.17-1.86 (m, 3H) ppm.

15 EXAMPLE 9

trans-(4a,9b)-2,5-dimethyl-6-[(4-methylphenyl)sulfanyl]-8-phenylsulfanyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole

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To an Et₂O solution (0.5 mL) of trans-(4a,9b)-8-bromo-2,5-dimethyl-6-[(4-methylphenyl)sulfanyl]-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole (Example 10, 100 mg, 0.25 mmol) and TMEDA (58 mg, 0.50 mmol) cooled in an acetone-dry ice bath was added dropwise t-BuLi (1.7 M pentane solution, 0.29 mL, 0.49 mmol). Resulted solution was allowed to maintain at -78 °C for 15 min at which time an Et₂O solution (0.5 mL) of S-phenylbenzosulfonate (124 mg, 0.49 mmol) was added dropwise over 2 min. Cooling bath was removed and reaction was allowed to warm

to rt over 20 min. This reaction was then allowed to stir at rt for additional 20 min before it was poured onto ice-cooled 1 M H₂SO₄ aq solution (5 mL). This mixture was extracted with CHCl₃ (3 x 10 mL), organic phases were combined, dried (MgSO₄) and concentrated. Resulted residue was then purified by SGC (5% MeOH/CH₂Cl₂) to provide the title compound (35 mg, 28%) as a light brown colored oil. ¹H NMR (CDCl₃, 300 MHz) δ 7.32 (d, 1H, J = 0.8 Hz), 7.24-7.01 (m, 10H), 3.37 (dd, 1H, J = 10.3, 3.0 Hz), 3.09 9s, 3H), 2.83 (t, 1H, J = 11.0 Hz), 2.60-2.51 (m, 1H), 2.40 (s, 3H), 2.29 (s, 3H), 2.20-2.01 (m, 2H), 1.89-1.77 (m, 1H) ppm.

10 EXAMPLE 10

di(*tert*-butyl) *cis*-(4a,9b)-8-bromo-6-methyl-3,4,4a,9b-tetrahydro-1H-pyrido[4,3-b]indole-2,5-dicarboxylate

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Step A. *o*-Tolylphenylhydrazine hydrogen chloride (25 g, 158.6 mmol) and 4-piperidone monohydrate hydrogen chloride (24.4 g, 158.6 mmol) were suspended in EtOH (310 mL). Concentrated aqueous HCl (26 g, 317 mmol)was then added. The mixture was refluxed for 3 h, and then the reaction was cooled to rt. The ppt was filtered and washed with cold EtOH. The white solid was air-dried for 18 h. 6-methyl-2,3,4,5-tetrahydro-1H-pyrido[4,3-b]indole hydrochloride (36.6 g, 141.2 mmol, 89%) was isolated and a white powder. ¹H NMR (CD₃OD, 300 MHz) δ 7.22-7.25 (1H, m), 6.89-6.96 (2H, m), 4.40 (2H, m), 3.60 (2H, t, 6.2 Hz), 3.16 (2H, t, 6.2 Hz), 2.45 (3H, s) ppm.

Step B. 6-Methyl-2,3,4,5-tetrahydro-1H-pyrido[4,3-b]indole hydrochloride (36.6 g, 141.2 mmol) was suspended in TFA (200 mL). The mixture was cooled to 0°C. Et₃SiH (32.86g, 282.6 mmol) was added slowly. The reaction was stirred at rt

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for 18 h. Add hexane (2x 300 mL), separate the acid layer, then basified with 50% NaOH until pH=14. Extract reaction with CHCl₃ (3 x 300 mL). The combined organic layers were washed with brine, dried, and concentrated to afford a light brown amorphous solid cis-6-methyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole (20.7 g, 110.1mmol, 78%). 1 H NMR (CDCl₃, 300 MHz) δ 6.90 (1H, d, 7.3 Hz), 6.82 (1H, d, 7.4 Hz), 6.62 (1H, t, 7.3 Hz), 3.77-3.82 (1H, m), 2.65-3.08 (5H, m), 2.13 (3H, s), 1.72-1.92 (2H, m) ppm.

Step C. *Cis*-(4a,9b)-6-methyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole (20.7g, 110.1 mmol) and BOC₂O (96.25g, 440.4 mmol) is heated at 110 °C for 3 hrs. Cool to rt and quench with water (20 mL). Add brine (200 mL) and EtOAc (200 mL); stir 10 min. Separate layers and re-extract aqueous with EtOAc (2 x 100 mL). Wash combined organic layers with brine, dry, and conc. 43 g of a brown, amorphous solid was isolated. The residue was purified by column chromatography (20-40% EtOAc/Hexane). Di(*tert*-butyl) *cis*-(4a,9b)-6-methyl-3,4,4a,9b-tetrahydro-1H-pyrido[4,3-b]indole-2,5-dicarboxylate (42 g, 108.2 mmol, 98%)was isolated as a clear viscous oil. ¹H NMR (DMSO, 400 MHz) δ 6.9-7.1 (3H, m), 4.5-4.7 (2H, m), 4.1-4.3 (1H, m), 3.55-3.65 (1H, m), 3.30-3.50 (2H, m), 2.8-3.0 (1H, m), 2.20 (3H, s), 1.50 (9H, s), 1.30 (9H, s) ppm.

Step D. Di(tert-butyl) cis-(4a,9b)-6-methyl-3,4,4a,9b-tetrahydro-1H-20 pyrido[4,3-b]indole-2,5-dicarboxylate (1,73g, 4.46 mmol) was dissolved in DMF (8 mL); the solution was then cooled to 0°C. NBS (0.874g, 4.90 mmol) was added as a solution in DMF(8 mL). The reaction was warmed to rt and stirred for 1 h. Add brine (10 mL) and EtOAc (10 mL). Stir 10 min and separate layers. Re-extract aqueous with EtOAc (2 x 20 mL). Wash combined organic with brine, dry, and concentrate. 25 1.88 g of an orange viscous oil was isolated. This crude residue was purified by column chromatography (10-30 % acetone/hexane). Di(tert-butyl) cis-(4a,9b)-8bromo-6-methyl-3,4,4a,9b-tetrahydro-1H-pyrido[4,3-b]indole-2,5-dicarboxylate (1.12g, 2.41 mmol, 54%) was isolated as a pale-yellow, amorphous solid. ¹H NMR (DMSO d₆, 400 MHz) δ 7.52 (1H, d, 7.5 Hz), 7.29-7.36 (2H, m), 4.55 (1H, ddd, 5.2 30 Hz, 6.4 Hz, 9.7 Hz), 3.66 (1H, dd, 3.7 Hz, 14.2 Hz), 3.64-3.66 (1H, m), 3.46 (1H, dd, 4.1 Hz, 13.9 Hz), 3.30-3.36 (1H, m), 2.00-3.09 (2H, m), 2.07-2.15 (1H, m), 1.65-1.76 (1H, m) ppm.

EXAMPLE 11

Di(tert-butyl) (4aS,9bR)-8-bromo-6-methyl-3,4,4a,9b-tetrahydro-1H-pyrido[4,3-b]indole-2,5-dicarboxylate

Step A. Di(*tert*-butyl)(4aS,9bR)-6-methyl-3,4,4a,9b-tetrahydro-1H-pyrido[4,3-b]indole-2,5-dicarboxylate was obtained from Di(*tert*-butyl) *cis*-(4a,9b)-6-methyl-3,4,4a,9b-tetrahydro-1H-pyrido[4,3-b]indole-2,5-dicarboxylate (Example 32 Step C)by using preparative HPLC on a ChiralPak® AD column (1% IPA in hexane).

Step B. Di(*tert*-butyl) (4aS,9bR)-6-methyl-3,4,4a,9b-tetrahydro-1H-pyrido[4,3-b]indole-2,5-dicarboxylate (20.2g, 51.99 mmol) was brominated according to the procedure of Example 13, Step D to give the title compound (21.3g, 45.7 mmol, 88%) as a pale-yellow, amorphous solid. ¹H NMR (DMSO, 400 MHz) identical to Example 13.

EXAMPLE 12

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di(*tert*-butyl) (4aS,9bR)-8-amino-6-methyl-3,4,4a,9b-tetrahydro-1H-pyrido[4,3-b]indole-2,5-dicarboxylate

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A solution of di(*tert*-butyl) (4aS,9bR)-8-bromo-6-methyl-3,4,4a,9b-tetrahydro-1H-pyrido[4,3-b]indole-2,5-dicarboxylate (Example 14) (9.35 g, 20.0 mmol), benzophenone imine (4.35g, 24.0 mmol), (S)-(-)-2,2'-bis(diphenylphosphino)-1,1'-binaphthyl (BINAP) (934 mg, 1.5 mmol), Pd2dba3 (1.04g, 1.0 mmol) and sodium *tert*-butoxide (2.7g, 28.0 mmol) in degassed toluene (325 mL) was heated at reflux under Argon atmosphere for 18 h. The solution was cooled and filtered through a pad of silica gel and eluted with EtOAc. The volatiles were removed under reduced pressure. The residue was taken up in methanol (200 mL) and then there was added NaOAC (3.43 g, 41.76 mmol) and hydroxylamine hydrochloride (2.2g, 31.32 mmol) and the mixture was stirred at rt for 1 h. The volatiles were removed under reduced pressure and the residue purified by column chromatography (20, 70% ethyl acetate/hexane) to afford the title compound (5.17g, 12.8mmol, 74%). MS (APCI): 404 (base, M+H).

General method for preparation of aryl-[(4aS,9bR)-6-methyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl]-amine

Step A. A solution of di(*tert*-butyl) (4aS,9bR)-8-amino-6-methyl-3,4,4a,9b-tetrahydro-1H-pyrido[4,3-b]indole-2,5-dicarboxylate (Example 18, 93.3 mg, 0.23 mmol), arylbromide (0.23mmol), BINAP (4.6 mg, 0.0069 mmol), Pd2dba3 (2.4 mg, 0.0023 mmol) and sodium tert-butoxide (44.5 mg, 0.46 mmol) in degassed toluene (2 mL) was heated at 80 °C for 18 h. The solution was cooled and filtered through a pad of silica gel and elute with EtOAc. The solvents were removed under reduced pressure and the residue purified by column chromatography (10, 20, 30% ethyl acetate/hexane) to afford di(tert-butyl) (4aS,9bR)-8-anilino-6-methyl-3,4,4a,9b-tetrahydro-1H-pyrido[4,3-b]indole-2,5-dicarboxylate.

Step B. A solution of di(*tert*-butyl) (4aS,9bR)-8-anilino-6-methyl-3,4,4a,9b-tetrahydro-1H-pyrido[4,3-b]indole-2,5-dicarboxylate in 20% TFA/CH₂Cl₂ (1 mL) was stirred at rt for 1 h. The reaction mixture was basified with NH₄OH to pH = 12, then extracted with CHCl₃ (3 x 3 mL). The combined organic layer was washed with brine, dried, and concentrated to afford the title compounds.

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(2,3-dichlorophenyl)-[(4aS,9bR)-6-methyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl]-amine

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The title compound (40 mg, 50 %) was prepared by the general method from di(*tert*-butyl) (4aS,9bR)-8-amino-6-methyl-3,4,4a,9b-tetrahydro-1H-pyrido[4,3-10 b]indole-2,5-dicarboxylate and 1-bromo-2,3-dichlorobenzene. ¹H NMR (CDCl₃, 300 MHz) δ 6.95 (1H, t, 8.1 Hz), 6.75-6.80 (4H, m), 6.02 (1H, s), 3.90-3.96 (1H, m), 3.53 (1H,bs), 2.74-3.11 (5H, m), 2.15 (3H, s), 1.65-1.87 (3H, m) ppm. MS (APCI): 348 (base, M+H).

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EXAMPLE 14

(3,4-dichlorophenyl)-[(4aS,9bR)-6-methyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl]-amine

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The title compound (27 mg, 34 %) was prepared by the general method from di(*tert*-butyl) (4aS,9bR)-8-amino-6-methyl-3,4,4a,9b-tetrahydro-1H-pyrido[4,3-b]indole-2,5-dicarboxylate and 4-bromo-1,2-dichlorobenzene. ¹H NMR (CDCl₃, 300 MHz) δ 7.17 (1H, d, 8.8 Hz), 6.85 (1H, d,2.6Hz), 6.70 -6.76 (2H, m), 6.61 (1H, dd, 8.8 Hz, 1.8 Hz), 5.44 (1H, s), 3.90-3.95 (1H, m), 3.53 (1H,bs), 2.73-3.09 (5H, m), 2.14 (3H, s), 1.67-1.98 (3H, m) ppm. MS (APCI): 348 (base, M+H).

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EXAMPLE 15

(3-chloro-4-methylphenyl)-[(4aS,9bR)-6-methyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl]-amine

The title compound (30 mg, 40 %) was prepared by the general method from di(*tert*-butyl) (4aS,9bR)-8-amino-6-methyl-3,4,4a,9b-tetrahydro-1H-pyrido[4,3-b]indole-2,5-dicarboxylate and 4-bromo-2-chloro-1-methylbenzene. ¹H NMR (CDCl₃, 300 MHz) δ 7.01 (1H, d, 8.1 Hz), 6.83 (1H, d, 2.2Hz), 6.75 (1H, s), 6.69 (1H, s), 6.64 (1H, dd, 8.4 Hz, 2.2 Hz), 5.34 (1H, s), 3.88-3.94 (1H, m), 3.53 (1H,bs), 2.73-3.12 (5H, m), 2.26 (3H, s), 2.13 (3H, s), 1.65-1.89 (3H, m) ppm. MS (APCI): 328 (base, M+H).

EXAMPLE 16

(2,4-dichlorophenyl)-[(4aS,9bR)-6-methyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl]-amine

The title compound (40 mg, 50 %) was prepared by the general method from di(*tert*-butyl) (4aS,9bR)-8-amino-6-methyl-3,4,4a,9b-tetrahydro-1H-pyrido[4,3-b]indole-2,5-dicarboxylate and 1-bromo-2,4-dichlorobenzene. ¹H NMR (CDCl₃, 300 MHz) δ 7.27 (1H, d, 2.6 Hz), 6.99 (1H, dd, 8.8 Hz, 2.6 Hz), 6.81 (2H, d, 8.8Hz), 6.75

(1H, s), 5.85 (1H, s), 3.90-3.95 (1H, m), 3.53 (1H,bs), 2.73-3.12 (5H, m), 2.14 (3H, s), 1.64-1.88 (3H, m) ppm. MS (APCI): 348 (base, M+H).

EXAMPLE 17

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(2,6-dichlorophenyl)-[(4aS,9bR)-6-methyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl]-amine

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The title compound (15 mg, 19 %) was prepared by the general method from di(tert-butyl) (4aS,9bR)-8-amino-6-methyl-3,4,4a,9b-tetrahydro-1H-pyrido[4,3-b]indole-2,5-dicarboxylate and 2-bromo-1,3-dichlorobenzene. ¹H NMR (CDCl₃, 300 MHz) δ 7.31 (2H, d, 8.0 Hz), 6.90-6.96 (1H, m), 6.46 (1H, s), 6.41 (1H, s), 5.76 (1H, s), 3.85-3.89 (1H, m), 3.53 (1H,bs), 2.72-3.06 (5H, m), 2.11 (3H, s), 1.65-1.91 (3H, m) ppm. MS (APCI): 348 (base, M+H).

EXAMPLE 18

20 (2,4-difluorophenyl)-[(4aS,9bR)-6-methyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl]-amine

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The title compound (11 mg, 15 %) was prepared by the general method from di(*tert*-butyl) (4aS,9bR)-8-amino-6-methyl-3,4,4a,9b-tetrahydro-1H-pyrido[4,3-b]indole-2,5-dicarboxylate and 1-bromo-2,4-dichlorobenzene. ¹H NMR (CDCl3, 300

MHz) δ 6.78-6.97 (3H, m), 6.77 (1H, s), 6.70 (1H, s), 5.37 (1H, s), 3.90-3.93 (1H, m), 3.53 (1H,bs), 2.75-3.10 (5H, m), 2.14 (3H, s), 1.68-1.88 (3H, m) ppm. MS (APCI): 316 (base, M+H).

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EXAMPLE 19

(2-methoxy-5-methylphenyl)-[(4aS,9bR)-6-methyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl]-amine

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The title compound (29 mg, 39 %) was prepared by the general method from di(*tert*-butyl) (4aS,9bR)-8-amino-6-methyl-3,4,4a,9b-tetrahydro-1H-pyrido[4,3-b]indole-2,5-dicarboxylate and 2-bromo-1-methoxy-4-methylbenzene. ¹H NMR (CDCl₃, 300 MHz) δ 6.71-6.84 (4H, m), 6.54 (1H, d, 0.7 Hz), 5.85 (1H, s), 3.91-3.96 (1H, m), 3.85 (1H, s), 3.53 (1H,bs), 2.80-3.17 (5H, m), 2.20 (3H, s), 2.15 (3H, s), 1.70-1.95 (2H, m) ppm. MS (APCI): 324 (base, M+H).

EXAMPLE 20

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2-chloro-4-[(4aS,9bR)-6-methyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-ylamino]-benzonitrile

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The title compound (50 mg, 64 %) was prepared by the general method from di(*tert*-butyl) (4aS,9bR)-8-amino-6-methyl-3,4,4a,9b-tetrahydro-1H-pyrido[4,3-

b]indole-2,5-dicarboxylate and 4-bromo-2-chloro-benzonitrile. ¹H NMR (CDCl₃, 300 MHz) δ 7.48 (1H, d, 8.5 Hz), 6.58-6.76 (4H, m), 5.11 (1H, s), 3.93-3.97 (1H, m), 3.61 (1H, bs), 2.76-3.14 (5H, m), 3.61 (1H,bs), 2.14 (3H, s), 1.70-1.90 (2H, m) ppm. MS (APCl): 339 (base, M+H).

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EXAMPLE 21

(4-chloro-2-fluorophenyl)-[(4aS,9bR)-6-methyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl]-amine

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The title compound (26 mg, 34 %) was prepared by the general method from di(*tert*-butyl) (4aS,9bR)-8-amino-6-methyl-3,4,4a,9b-tetrahydro-1H-pyrido[4,3-b]indole-2,5-dicarboxylate and 1-bromo-4-chloro-2-fluorobenzene. ¹H NMR (CDCl₃, 300 MHz) δ 7.01-7.05 (1H, m), 6.84-6.92 (2H, m), 6.73-6.79 (2H, m), 5.53 (1H, s), 3.89-3.93 (1H, m), 3.53 (1H,bs), 2.75-3.11 (5H, m), 2.14 (3H, s), 1.68-1.89 (3H, m) ppm. MS (APCI): 332 (base, M+H).

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EXAMPLE 22

(2-fluoro-5-trifluoromethyl-phenyl)-[(4aS,9bR)-6-methyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl]-amine

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The title compound (37 mg, 44 %) was prepared by the general method from di(*tert*-butyl) (4aS,9bR)-8-amino-6-methyl-3,4,4a,9b-tetrahydro-1H-pyrido[4,3-b]indole-2,5-dicarboxylate and 2-bromo-1-fluoro-4-trifluoromethyl-benzene. ¹H NMR (CDCl₃, 300 MHz) δ 7.04-7.15 (2H, m), 6.89-6.94 (1H, m), 6.82 (1H, s), 6.77 (1H, s), 5.72 (1H, s), 3.92-3.97 (1H, m), 3.59 (1H,bs), 2.75-3.12 (5H, m), 2.16 (3H, s), 1.66-1.87 (3H, m) ppm. MS (APCI): 366 (base, M+H).

EXAMPLE 23

10 (2-chloro-5-trifluoromethyl-phenyl)-[(4aS,9bR)-6-methyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl]-amine

The title compound (37 mg, 42 %) was prepared by the general method from di(*tert*-butyl) (4aS,9bR)-8-amino-6-methyl-3,4,4a,9b-tetrahydro-1H-pyrido[4,3-b]indole-2,5-dicarboxylate and 2-bromo-1-chloro-4-trifluoromethyl-benzene. ¹H NMR (CDCl₃, 300 MHz) δ 7.35-7.38 (1H, m), 7.06 (1H, d, 1.9 Hz), 6.86-6.90 (1H, m), 6.74-6.83 (2H, m), 6.04 (1H, s), 3.95-3.97 (1H, m), 3.59 (1H,bs), 2.76-3.14 (5H, m), 2.16 (3H, s), 1.72-1.95 (3H, m) ppm. MS (APCI): 382 (base, M+H).

cis-(4a,9b)-8-bromo-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid tert-butyl ester

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Step A. To a solution of (2-trifluoromethyl-phenyl)-hydrazine (12.0 g, 68.13 mmol) and 4-piperidone monohydrate (10.46 g, 68.09 mmol) in *iso*-propyl alcohol was bubbled HCl (g) for 15 minutes at rt. The reaction was sealed in a pressure vessel and heated to 95 °C for 5 h, then cooled to rt. The reaction mixture was filtered and washed with iso-propyl alcohol and dried in vacuo to yield 6-trifluoromethyl-2,3,4,5-tetrahydro-1H-pyrido[4,3-b]indole bis-hydrochloride as a white solid (13.3 g, 63% yield). ¹H NMR (CD₃OD, 300 MHz) δ 3.203 (t, 2H, 12.4 Hz), 3.642 (t, 2H, 12.0 Hz), 4.472 (t, 2H, 2.6 Hz), 7.189 (t, 1H, 15.7 Hz), 7.435 (d, 1H, 7.7 Hz), 7.709 (d, 1H, 8.0 Hz) ppm.

Step B. A solution of 6-trifluoromethyl-2,3,4,5-tetrahydro-1H-pyrido[4,3-b]indole bis-hydrochloride (13.2 g, 42.15 mmol) in trifluoroacetic acid was stirred for 30 min at rt to dissolve the starting material. Triethyl-silane (9.88 g, 84.94 mmol) was added to form a biphasic mixture. The reaction was let stir at rt for 90 h. The reaction mixture was concentrated in vacuo. The reaction was then neutralized with base and extracted with chloroform. The organic layer was concentrated in vacuo to yield *cis*-(4a,9b)-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole as an off white solid (11.3 g). ¹H NMR (CDCl₃, 300 MHz) δ 1.609-1.707 (m, 1H), 1.739 (s, 1H), 1.806-1.912 (m, 1H), 2.713-2.789 (m, 1H), 2.859-2.985 (m, 2H), 3.026-3.131 (m, 2H), 3.970 (dd, 1H, 5.1, 10.3 Hz), 4.308 (s, 1H), 6.755 (t, 1H, 15.4 Hz), 7.190-7.243 (m, 2H) ppm.

Step C. To a solution of crude cis-(4a,9b)-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole (11.3 g, < 46.65 mmol) in 1,4-dioxane was added

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1M NaOH in water (93.3 mL, 93.3 mmol) and let stir 5 min at rt. To the reaction mixture was added dropwise a solution of (Boc)₂O (12.2 g, 55.98 mmol) in 1,4-dioxane and let stir 18 h at rt. The reaction mixture was then quenched with water, washed with acid and extracted with ether. The organic layer was concentrated in vacuo and triturated with hexane and filtered to obtain *cis*-(4a,9b)-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester as an off white solid (10.6 g, 30.96 mmol). ¹H NMR (CDCl₃, 300 MHz) δ 1.457 (s, 9H), 1,525 (s, 1H), 1.760-1.820 (m, 1H), 1.840-2.020 (m, 1H), 2.280-2.400 (m, 2H), 2.410-2.580 (m, 2H), 2.720-2.840 (m, 1H), 4.080-4.180 (m, 1H), 6.740-6.800 (t, 1H, 18.0 Hz), 7.238 (s, 1H), 7.262 (s, 1H) ppm.

Step D. To *cis*-(4a,9b)-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (1.0 g, 2.92 mmol) in DMF at 0 °C was added dropwise solution of NBS (520 mg, 2.92 mmol) in DMF and let stir for 30 min at 0°C. The reaction mixture was then warmed to rt and let stir for 60 min. The reaction mixture was quenched with water and extracted with ether. The organic layer was concentrated in vacuo and triturated with hexane and filtered to obtain the title compound as a yellow solid (902 mg, 73% yield). 1 H NMR (CDCl3, 300 MHz) 5 1.471 (s, 9H), 1.650-1.800 (m, 1H), 1.880-2.000 (m, 1H), 3.300-3.400 (m, 2H), 3.420-3.580 (m, 2H), 3.620-3.780 (m, 2H), 4.080-4.180 (m, 1H), 7.340 (s, 1H), 7.357 (s, 1H) ppm.

EXAMPLE 25

(4aS,9bR)-8-bromo-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester

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Step A. (4aS,9bR)-6-Trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid tert-butyl ester was obtained from *cis*-(4a,9b)-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 30 step C) by using preparative HPLC on a ChiralPak® AD column (2% IPA in hexane).

Step B. The title compound was prepared by following the bromination method as exemplified by the step D of Example 30. ¹H NMR (DMSO d₆, 400 MHz) identical to Example 30.

General method for preparation of alkyl(or benzyl)-[cis-(4a,9b)-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1*H*-pyrido[4,3-*b*]indol-8-yl]-amine

Step A. To a solution of *cis*-(4a,9b)-8-bromo-6-trifluomethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-*b*]indole-2-carboxylic acid *tert*-butyl ester (1.0 mole equivalent) in toluene was added alkyl(or benzyl) amines (3.0 mole equivalent),

NaOt-Bu (3.0 mole equivalent), (O-biPh)P(tBu)2 (0.18 mole equivalent) and

Pd2(dba)3 (0.06 mole equivalent). The reaction mixture was heated at 80°C for 16-20 h, then cooled to rt. The reaction mixture was then quenched with water, washed with base and extracted with ethyl acetate. The organic layer was concentrated *in vacuo* and chromatographed on a silica gel column by elution with Hexane/Ethyl Acetate to give *cis*-(4a,9b)-8-alkyl(or benzyl)amino-6-trifluoromethyl-1-2,3,4,4a,5,9b-hexahydro-pyrido[4,3-*b*]indole-2-carboxylic acid *tert*-butyl ester.

Step B. A solution of *cis*-(4a,9b)-8-alkyl(or benzyl)amino-6-trifluoromethyl-1-2,3,4,4a,5,9b-hexahydro-pyrido[4,3-*b*]indole-2-carboxylic acid *tert*-butyl ester in 20% trifluoroacetic acid in dichloromethane was stirred for 1-2 h at rt. The reaction mixture was then concentrated *in vacuo* and then neutralized with base and extracted with chloroform. The organic layer was concentrated *in vacuo* to give the title compound (12-68% overall yield).

cyclohexyl-[cis-(4a,9b)-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl]-amine

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The title compound was prepared by following the general method as a yellow solid (15 mg, 44%) from *cis*-(4a,9b)-8-bromo-6-trifluomethyl-1,3,4,4a,5,9b
hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (42 mg, 0.10 mmol) and cyclohexylamine (30 mg, 0.30 mmol). ¹H NMR (CDCl3, 300 MHz) δ 1.054
1.383 (m, 6H), 1.630-1.774 (m, 4H), 1.849-1.910 (m, 1H), 1.994-2.038 (m, 2H), 2.763-3.190 (m, 6H), 3.890-3.908 (m, 2H), 6.477 (s, 1H), 6.594 (s, 1H) ppm.

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EXAMPLE 27

(1,3-Dimethyl-butyl)-[cis-(4a,9b)-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl]-amine

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The title compound was prepared by following the general method as a yellow oil (4 mg, 12%) from cis-(4a,9b)-8-bromo-6-trifluomethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid tert-butyl ester (42 mg, 0.10 mmol) and 1,3-dimethyl-butylamine (30 mg, 0.30 mmol). 1 H NMR (CDCl₃, 300 MHz) δ 0.835-0.900 (m, 9H), 1.060 (d, 3H, 6.2 Hz), 1.153-1.184 (m, 4H), 1.480-1.850 (m, 1H),

2.780-3.190 (m, 4H), 3.280-3.480 (m, 1H), 3.820-3.980 (m, 2H), 6.413 (s, 1H), 6.514 (s, 1H) ppm.

EXAMPLE 28

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Benzyl-[cis-(4a,9b)-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl]-amine

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The title compound was prepared by following the general method as a yellow solid (24 mg, 68%) from cis-(4a,9b)-8-bromo-6-trifluomethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid tert-butyl ester (42 mg, 0.10 mmol) and benzylamine (32 mg, 0.30 mmol). 1 H NMR (CDCl₃, 300 MHz) δ 1.697-1.708 (m, 1H), 1.855-1.900 (m, 1H), 2.192 (s, 2H), 2.772-2.819 (m, 1H), 2.896-3.108 (m, 4H), 3.911-3.963 (m, 2H), 4.276 (s, 2H), 6.559 (s, 1H), 6.661 (s, 1H), 7.300-7.384 (m, 5H) ppm.

EXAMPLE 29

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(1-phenyl-ethyl)-[cis-(4a,9b)-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl]-amine

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The title compound was prepared by following the general method as a yellow oil (8 mg, 22%) from cis-(4a,9b)-8-bromo-6-trifluomethyl-1,3,4,4a,5,9b-hexahydropyrido[4,3-b]indole-2-carboxylic acid tert-butyl ester (42 mg, 0.10 mmol) and 1-phenyl-ethylamine (36 mg, 0.30 mmol). 1 H NMR (CDCl₃, 300 MHz) δ 1.253 (s, 1H), 1.481-1.508 (m, 3H), 1.618 (d, 1H, 6.9 Hz), 1.807-1.899 (m, 4H), 2.748-2.838 (m, 1H), 2.928-3.034 (m, 3H), 3.80-3.92 (m, 2H), 6.404 (d, 1H, 14.0 Hz), 6.512 (d, 1H, 9.5 Hz), 7.287-7.393 (m, 5H) ppm.

EXAMPLE 30

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(2-methyl-benzyl)-[cis-(4a,9b)-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl]-amine

H H NH F H

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The title compound was prepared by following the general method as a yellow solid (20 mg, 55%) from cis-(4a,9b)-8-bromo-6-trifluomethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid tert-butyl ester (42 mg, 0.10 mmol) and 2-methyl-benzylamine (36 mg, 0.30 mmol). 1H NMR (CDCl₃, 300 MHz) δ 1.717-1.779 (m, 1H), 1.935-2.015 (m, 1H), 2.377 (s, 3H), 2.476-2.680 (m, 2H), 2.827-2.922 (m, 2H), 2.999-3.194 (m, 3H), 3.918-3.967 (m, 2H), 4.199 (s, 2H), 6.548 (s, 1H), 6.632 (s, 1H), 7.137-7.244 (m, 3H), 7.292-7.317 (m, 1H) ppm.

(2-methoxybenzyl)-[cis-(4a,9b)-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl]-amine

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The title compound was prepared by following the general method as a yellow solid (20 mg, 53%) from *cis*-(4a,9b)-8-bromo-6-trifluomethyl-1,3,4,4a,5,9b-10 hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (42 mg, 0.10 mmol) and 2-methoxy-benzylamine (41 mg, 0.30 mmol). ¹H NMR (CDCl₃, 300 MHz) δ 1.708-1.719 (m, 1H), 1.911-1.927 (m, 1H), 2.400-2.600 (m, 2H), 2.808-2.903 (m, 2H), 2.960-3.122 (m, 3H), 3.864 (s, 3H), 3.879-3.933 (m, 2H), 4.255 (s, 2H), 6.579 (s, 1H), 6.657 (s, 1H), 6.879-6.941 (m, 3H), 7.278-7.290 (m, 1H) ppm.

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EXAMPLE 32

(2-chloro-6-fluoro-benzyl)-[cis-(4a,9b)-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl]-amine

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The title compound was prepared by following the general method as an orange solid (13 mg, 34%) from *cis*-(4a,9b)-8-bromo-6-trifluomethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (42 mg, 0.10 mmol)

and 2-chloro-6-fluoro-benzylamine (48 mg, 0.30 mmol). ¹H NMR (CDCl₃, 300 MHz) δ 1.796-1.808 (m, 1H), 2.015-2.028 (m, 1H), 2.220-2.500 (m, 5H), 2.828-3.215 (m, 4H), 4.010 (d, 1H, 1.8 Hz), 4.445 (s, 1H), 6.679 (s, 1H), 6.758 (s, 1H), 6.974-7.180 (m, 1H), 7.190-7.233 (m, 2H) ppm.

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EXAMPLE 33

(4-tert-butyl-benzyl)-[cis-(4a,9b)-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl]-amine

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The title compound was prepared by following the general method as an orange solid (16 mg, 39%) from *cis*-(4a,9b)-8-bromo-6-trifluomethyl-1,3,4,4a,5,9b-15 hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (42 mg, 0.10 mmol) and 4-*tert*-butyl-benzylamine (49 mg, 0.30 mmol). ¹H NMR (CDCl₃, 300 MHz) δ 1.342 (s, 9H), 1.720-1.820 (m, 1H), 1.950-2.050 (m, 1H), 2.550-2.650 (m, 2H), 2.820-2.980 (m, 2H), 3.020-3.250 (m, 3H), 3.880-4.000 (m, 2H), 4.229 (s, 2H), 6.576 (s, 1H), 6.660 (s, 1H), 7.357 (dd, 4H, 8.5, 26.0 Hz) ppm.

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(3-methylbenzyl)-[cis-(4a,9b)-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl]-amine

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The title compound was prepared by following the general method as a yellow solid (14 mg, 38%) from *cis*-(4a,9b)-8-bromo-6-trifluomethyl-1,3,4,4a,5,9b-10 hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (42 mg, 0.10 mmol) and 3-methyl-benzylamine (36 mg, 0.30 mmol). ¹H NMR (CDCl₃, 300 MHz) δ 1.695-1.756 (m, 1H), 1.921-1.966 (m, 1H), 2.354 (s, 3H), 2.382-2.441 (m, 2H), 2.829-2.885 (m, 2H), 2.979-3.157 (m, 3H), 3.908-3.955 (m, 2H), 4.207 (s, 2H), 6.551 (s, 1H), 6.642 (s, 1H), 7.086-7.239 (m, 4H) ppm.

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EXAMPLE 35

(4-methylbenzyl)-[cis-(4a,9b)-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl]-amine

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The title compound was prepared by following the general method as a yellow solid (16 mg, 45%) from *cis*-(4a,9b)-8-bromo-6-trifluomethyl-1,3,4,4a,5,9b-

25 hexahydro-pyrido[4,3-b]indole-2-carboxylic acid tert-butyl ester (42 mg, 0.10 mmol)

and 4-methyl-benzylamine (36 mg, 0.30 mmol). ¹H NMR (CDCl₃, 300 MHz) δ 1.698-1.759 (m, 1H), 1.925-1.972 (m, 1H), 2.366 (s, 3H), 2.400-2.464 (m, 2H), 2.831-2.903 (m, 2H), 2.988-3.163 (m, 3H), 3.920-3.967 (m, 2H), 4.223 (s, 2H), 6.563 (s, 1H), 6.654 (s, 1H), 7.161-7.188 (d, 2H, 8.0 Hz), 7.225 (s, 2H) ppm.

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EXAMPLE 36

(2,5-Dimethylbenzyl)-[cis-(4a,9b)-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl]-amine

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The title compound was prepared by following the general method as a yellow oil (17 mg, 46%) from cis-(4a,9b)-8-bromo-6-trifluomethyl-1,3,4,4a,5,9b-hexahydropyrido[4,3-b]indole-2-carboxylic acid tert-butyl ester (42 mg, 0.10 mmol) and 2,5-dimethyl-benzylamine (41 mg, 0.30 mmol). 1 H NMR (CDCl₃, 300 MHz) δ 1.943-2.004 (m, 1H), 2.165-2.253 (m, 2H), 2.323 (s, 3H), 2.228 (s, 3H), 2.780 (dd, 1H, 10.2, 12.8 Hz), 3.155-3.360 (m, 4H), 3.990-4.033 (m, 3H), 4.162 (s, 2H), 6.595 (s, 1H), 6.650 (s, 1H), 7.034-7.175 (m, 3H) ppm.

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(3-fluoro-5-trifluoromethyl-benzyl)-[cis-(4a,9b)-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl]-amine

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The title compound was prepared by following the general method as a yellow oil (25 mg, 58%) from cis-(4a,9b)-8-bromo-6-trifluomethyl-1,3,4,4a,5,9b-hexahydropyrido[4,3-b]indole-2-carboxylic acid tert-butyl ester (42 mg, 0.10 mmol) and 3-fluoro-5-trifluoromethyl-benzylamine (58 mg, 0.30 mmol). ¹H NMR (CDCl₃, 300 MHz) δ 1.667-1.762 (m, 1H), 1.872-1.964 (m, 1H), 2.359-2.447 (m, 2H), 2.793-2.918 (m, 2H), 2.968-3.155 (m, 3H), 3.925-4.002 (m, 2H), 4.356 (s, 2H), 6.492 (s, 1H), 6.633 (s, 1H), 7.238-7.319 (m, 2H), 7.453 (s, 1H) ppm.

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EXAMPLE 38

cis-(4a,9b)-8-bromo-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydropyrido[4,3-b]indole-2-carboxylic acid tert-butyl ester

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Step A. To a solution of *cis*-(4a,9b)-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 30 step C, 1.03 g, 3.0 mmol, 1.0 mole equivalent) in DMF (15 mL) was added NaH (0.24 g, 10

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mmol, 3.3 mole equivalent). The reaction mixture was stirred at rt for 10 min before MeI (2 mL, 31 mmol, 10 mole equivalent) was added dropwise. The reaction mixture was stirred at rt for 0.5 h, quenched with water (100 mL) and extracted with ether (3×50 mL). The organic solution was dried (Na₂SO₄), concentrated in vacuo and the residue was chromatographed on a silica gel column by elution with EtOAc/Hexane (gradient) to give *cis*-(4a,9b)-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester as a white solid (0.71 g, 66%). ¹H NMR (CDCl₃, 300 MHz) δ (ppm) 1.41 (s, 9H), 1.78-2.12 (m, 2H), 2.94-3.00 (m, 3H), 3.22-3.58 (m, 4H), 3.60-3.82 (m, 2H), 6.67 (t, J = 7.2 Hz, 1H), 7.16 (d, J = 7.2 Hz, 1H).

Step B. To a solution *cis*-(4a,9b)-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (0.39 g, 1.1 mmol, 1.0 mole equivalent) in DMF (4 mL) was added NBS (0.20 g, 1.1 mmol, 1.0 mole equivalent) dropwise at 0 °C. The reaction mixture was stirred at 0 °C for 10 min, rt for 30 min, quenched with water (20 mL) and extracted with ether (3×10 mL). The ether solution was dried (Na₂SO₄), concentrated *in vacuo* and the residue was chromatographed on a silica gel column by elution with EtOAc/Hexane (gradient) to give the title compound as a yellow solid (0.40 g, 84%). ¹H NMR (CDCl₃, 300 MHz) δ (ppm) 1.42 (s, 9H), 1.70-2.12 (m, 2H), 2.90-2.98 (m, 3H), 3.15-3.80 (m, 6H), 7.22 (d, J = 1.7 Hz, 1H), 7.41(d, J = 1.7 Hz, 1H).

EXAMPLE 39

(4aS,9bR)-8-bromo-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydropyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester

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(4aS,9bR)-6-Trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 31Step A) was methylated and brominated according to the procedure of Example 44, Step A and B to give the title compound as a pale-yellow, amorphous solid. ¹H NMR (DMSO d₆, 400 MHz) identical to example 86.

General method for preparation of alkyl(or benzyl)-(5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1*H*-pyrido[4,3-b]indol-8-yl)-amine

Step A. A solution of 8-bromo-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b
hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 44 or 45,

1.0 mole equivalent), alkyl (or benzyl)amine (2.0 mole equivalent), NaOt-Bu (3.0

mole equivalent) in toluene was degassed at 50 °C under Ar for 10 min and then

cooled down to rt. A solution of Pd2(dba)3 (0.05 mole equivalent) and (o-biPh)P(t
Bu)2 (0.15 mole equivalent) in toluene was degassed at rt under Ar for 10 min and

added to the reaction mixture. The reaction mixture was then degassed at 50 °C under

Ar for 10 min, heated at 80 °C for 16 h, cooled to rt and quenched with water.

Hexane was added and the organic layer was loaded directly on a silica gel column.

Gradient elution with EtOAc/Hexane gave the 8-alkyl(or benzyl)amino-5-methyl-6
trifluoromethyl-1-2,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*
butyl ester.

Step B. To a solution of 8-alkyl(or benzyl)amino-5-methyl-6-trifluoromethyl-1-2,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (~0.1 mmol) in CH₂Cl₂ (4 mL) was added TFA (1 mL). The reaction mixture was stirred at rt for 2 h, concentrated *in vacuo*, basified with NH₄OH (10 mL) and extracted with CH₂Cl₂ (3×10 mL). The organic solution was dried (MgSO₄) and concentrated *in vacuo* to give the products in 35-75 % overall yields.

cyclohexyl-[cis-(4a,9b)-5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl]-amine

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The title compound was prepared by following the general method as a yellow solid (19 mg, 55%) from *cis*-(4a,9b)-8-bromo-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 44, 44 mg, 0.10 mmol) and cyclohexylamine (20 mg, 0.20 mmol). ¹H NMR (CDCl₃, 300 MHz) δ (ppm) 0.95-1.40 (m, 5H), 1.42-1.75 (m, 3H), 1.75-2.03 (m, 4H), 2.43 (dd, J = 12.8, 9.5 Hz, 1H), 2.74 (d, J = 1.5 Hz, 3H), 2.82-2.95 (m, 2H), 2.95-3.20 (m, 3H), 3.20-3.30 (m, 1H), 3.30-4.20 (m, 2H), 6.50 (s, 1H), 6.54 (d, J = 1.5 Hz, 1H).

EXAMPLE 41

benzyl-[cis-(4a,9b)-5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl]-amine

The title compound was prepared by following the general method as a yellow solid (23 mg, 64 %) from *cis*-(4a,9b)-8-bromo-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester

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(Example 44, 44 mg, 0.10 mmol) and benzylamine (21 mg, 0.20 mmol). 1 H NMR (CDCl₃, 300 MHz) δ (ppm) 1.87-2.03 (m, 2H), 2.64 (dd, J = 9.5, 12.8 Hz, 1H), 2.75-2.88 (m, 3H), 2.88-3.10 (m, 1H), 3.08 (dd, J = 6.2, 12.8 Hz, 1H), 3.23 (td, J = 6.2, 9.2 Hz, 1H), 3.32-3.42 (m, 1H), 3.75-3.85 (br, 3H), 4.27 (s, 2H), 6.62 (d, J = 2.2 Hz, 1H), 6.70 (d, J = 2.2 Hz, 1H), 7.20-7.42 (m, 5H).

EXAMPLE 42

[cis-(4a,9b)-5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl]-(2-trifluoromethyl-benzyl)-amine

The title compound was prepared by following the general method as a yellow solid (33 mg, 75 %) from *cis*-(4a,9b)-8-bromo-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 44, 44 mg, 0.10 mmol) and 2-trifluoromethyl-benzylamine (35 mg, 0.20 mmol). ¹H NMR (CDCl₃, 300 MHz) δ (ppm) 1.78-2.00 (m, 2H), 2.64 (dd, J = 9.5, 12.8 Hz, 1H), 2.75-2.95 (m, 4H), 3.02 (dd, J = 6.2, 12.8 Hz, 1H), 3.08-3.30 (m, 3H), 3.37 (td, J = 4.4, 8.6 Hz, 1H), 3.85-4.05 (br, 1H), 4.49 (s, 2H), 6.58 (d, J = 2.2 Hz, 1H), 6.63 (d, J = 2.2 Hz, 1H), 7.39 (t, J = 7.5 Hz, 1H), 7.52 (t, J = 7.5 Hz, 1H), 7.63 (d, J = 7.7 Hz, 1H), 7.70 (d, J = 7.7 Hz, 1H); ¹⁹F NMR (CDCl₃, 300 MHz) δ (ppm) - 55.14, -60.45.

[cis-(4a,9b)-5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl]-(1-phenyl-ethyl)-amine

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The title compound was prepared by following the general coupling procedure as a yellow solid (26 mg, 70 %) from *cis*-(4a,9b)-8-bromo-5-methyl-6trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 44, 44 mg, 0.10 mmol) and α-methylbenzylamine (24 mg, 0.20 mmol). ¹H NMR (CDCl₃, 300 MHz) δ (ppm) 1.51 (dd, J = 1.8, 6.6 Hz, 3H), 1.75-1.92 (m, 2H), 2.20-2.55 (br, 2H), 2.55-2.66 (m, 1H), 2.72-3.12 (m, 6H), 3.23-3.40 (m, 1H), 3.60-4.10 (br, 1H), 4.41 (q, J = 6.3 Hz, 1H), 6.48 (dd, J = 2.2, 5.5 Hz, 1H), 6.56 (dd, J = 2.6, 8.4 Hz, 1H), 7.18-7.42 (m, 5H).

EXAMPLE 44

((S)-1-cyclohexyl-ethyl)-[cis-(4a,9b)-5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl]-amine

The title compound was prepared by following the general method as a yellow solid (16 mg, 42 %) from *cis*-8-bromo-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 44, 44 mg,

0.10 mmol) and (S)-cyclohexyl-ethylamine (25 mg, 0.20 mmol). 1 H NMR (CDCl₃, 300 MHz) δ (ppm) 0.95-1.49 (m, 10H), 1.55-1.95 (m, 7H), 2.00-2.42 (br, 2H), 2.60-2.78 (m, 1H), 2.78-2.95 (m, 4H), 3.02 (dd, J = 6.2, 12.8 Hz, 1H), 3.08-3.30 (m, 2H), 3.30-4.20 (m, 1H), 6.55 (s, 1H), 6.58 (d, J = 2.4 Hz, 1H).

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EXAMPLE 45

(exo-bicyclo[2.2.1]hept-2-yl)-[cis-(4a,9b)-5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl]-amine

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The title compound was prepared by following the general procedure as a yellow solid (23 mg, 64 %) from *cis*-(4a,9b)-8-bromo-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 44, 44 mg, 0.10 mmol) and *exo*-2-aminonorborane (22 mg, 0.20 mmol).

¹H NMR (CDCl₃, 300 MHz) δ (ppm) 0.95-1.25 (m, 4H), 1.25-1.55 (m, 3H), 1.72 (dd, J = 7.7, 12.4 Hz, 1H), 1.78-1.88 (m, 2H), 2.08-2.22 (m, 2H), 2.50-2.65 (m, 1H), 2.74 (d, J = 1.8 Hz, 3H), 2.78-2.95 (m, 2H), 3.02 (dd, J = 6.2, 12.8 Hz, 1H), 3.05-3.35 (m, 20 5H), 6.47 (d, J = 2.4 Hz, 1H), 6.50 (d, J = 2.4 Hz, 1H).

EXAMPLE 46

[cis-(4a,9b)-5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl]-((S)-2-phenyl-propyl)-amine

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The title compound was prepared by following the general method as a yellow solid (13 mg, 34 %) from cis-(4a,9b)-8-bromo-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid tert-butyl ester (Example 44, 44 mg, 0.10 mmol) and (S)-2-phenyl-propanylamine (27 mg, 0.20 mmol). ¹H NMR (CDCl₃, 300 MHz) δ (ppm) 1.27 (d, J = 6.9 Hz, 3H), 1.98-2.18 (m, 1H), 2.40-2.55 (m, 1H), 2.73 (d, J = 1.5 Hz, 3H), 2.80-3.43 (m, 11H), 6.45 (s, 1H), 6.53 (d, J = 1.5 Hz, 1H), 7.00-7.32 (m, 5H).

General method for preparation of aryl-[(4aS,9bR)6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl]-amine

To a solution of (4aS,9bR)-8-bromo-5-methyl-6-trifluomethyl-1,3,4,4a,5,9bhexahydro-pyrido[4,3-b]indole-2-carboxylic acid tert-butyl ester (Example 45, 1.0 mole equivalent) in toluene was added substituted anilines (3.0 mole equivalent), NaOt-Bu (3.0 mole equivalent), BINAP (0.18 mole equivalent) and Pd2(dba)3 (0.06 mole equivalent). The reaction mixture was heated at 80°C for 16-20 h, then cooled to rt. The reaction mixture was then quenched with water, washed with base and extracted with ethyl acetate. The organic layer was concentrated in vacuo and chromatographed on a silica gel column by elution with Hexane/Ethyl Acetate to give 8-arylamino-5-methyl-6-trifluoromethyl-1-2,3,4,4a,5,9b-hexahydro-pyrido[4,3b]indole-2-carboxylic acid tert-butyl ester. A solution of 8-arylamino-5-methyl-6trifluoromethyl-1-2,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid tertbutyl ester in 20% trifluoroacetic acid in dichloromethane was stirred for 1-2 h at rt. The reaction mixture was then concentrated in vacuo and then neutralized with base and extracted with chloroform. The organic layer was concentrated in vacuo to give aryl-(5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8yl)-amine (12-68% yield).

(2-methylthio-phenyl)-[(4aS,9bR)-5-methyl-6-trifloromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl)]-amine

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The title compound was prepared by following the general coupling procedure as a yellow solid (21 mg, 54%) from (4aS,9bR)-8-bromo-5-methyl-6-trifluomethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 45, 41 mg, 0.10 mmol) and 2-methylthio-phenylamine (42 mg, 0.30 mmol). ¹H NMR (CDCl₃, 300 MHz) δ 1.896-1.980 (m, 2H), 2.386 (s,1H), 2.723-2.819 (m, 4H), 2.912-2.927 (m, 3H), 3.065-6.262 (m, 2H), 3.491-3.546 (m, 1H), 6.471 (s, 1H), 6.775 (dt, 1H, 1.1, 7.3 Hz), 6.908 (dd, 1H, 1.1, 8.4 Hz), 7.057 (d, 1H, 1.9 Hz), 7.121 (t, 1H, 6.7 Hz), 7.199 (d, 1H, 2.2 Hz), 7.429 (dd, 1H, 1.5, 7.7 Hz) ppm.

EXAMPLE 48

(2-ethylphenyl)-[(4aS,9bR)-5-methyl-6-trifloromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl)-amine

The title compound was prepared by following the general coupling procedure as a yellow solid (16 mg, 43%) from (4aS,9bR)-8-bromo-5-methyl-6-trifluomethyl-

1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 45, 41 mg, 0.10 mmol) and 2-ethyl-phenylamine (36 mg, 0.30 mmol). ¹H NMR (CDCl₃, 300 MHz) δ 1.274 (t, 3H, 7.4 Hz), 1.883-1.933 (m, 2H), 2.374 (s, 2H), 2.598 (dd, 2H, 7.7, 15.1 Hz), 2.678-2.747 (m, 1H), 2.820-2.979 (m, 4H), 3.041 (dd, 1H, 5.9, 12.8 Hz), 3.177 (dd, 1H, 6.6, 14.3 Hz), 3.457-3.509 (m, 1H), 5.261 (s, 1H), 6.883 (dt, 1H, 1.1, 7.3 Hz), 6.950-6.968 (m, 2H), 7.052-7.121 (m, 2H), 7.179 (d, 1H, 7.7 Hz) ppm.

EXAMPLE 49

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(2-methoxyl-5-methylphenyl)-[(4aS,9bR)-5-methyl-6-trifloromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl)-amine

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The title compound was prepared by following the general coupling procedure as a yellow solid (25 mg, 64%) from (4aS,9bR)-8-bromo-5-methyl-6-trifluomethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 45, 41 mg, 0.10 mmol) and 2-methoxy-5-methyl-phenylamine (41 mg, 0.30 mmol). ¹H NMR (CDCl₃, 300 MHz) δ 1.879-1.938 (m, 2H), 2.221 (s, 3H), 2.507 (s, 2H), 2.670-2.741 (m, 1H), 2.829-2.963 (m, 4H), 3.032 (dd, 1H, 5.9, 12.9 Hz), 3.197 (dd, 1H, 6.6, 14.3 Hz), 3.474-3.527 (m, 1H), 5.889 (s, 1H), 6.580 (dd, 1H, 1.1, 8.1 Hz), 6.731-6.779 (m, 2H), 7.059 (d, 1H, 1.8 Hz), 7.184 (d, 1H, 2.2 Hz) ppm.

[(4aS,9bR)-5-methyl-6-trifloromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl]-phenyl-amine

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The title compound was prepared by following the general coupling procedure as a yellow solid (25 mg, 90%) from (4aS,9bR)-8-bromo-5-methyl-6-trifluomethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 45, 35 mg, 0.08 mmol) and phenylamine (22 mg, 0.24 mmol). ¹H NMR (CDCl₃, 300 MHz) δ 1.945-2.068 (m, 2H), 2.693 (dd, 1H, 9.2, 12.9 Hz), 2.915 (d, 2H, 2.2 Hz), 2.950-3.022 (m, 2H), 3.135 (dd, 1H, 6.3, 12.9 Hz), 3.281-3.354 (m, 1H), 3.471-3.572 (m, 3H), 5.539 (s, 1H), 6.852-6.987 (m, 3H), 7.043 (d, 1H, 1.8 Hz), 7.172 (d, 1H, 2.2 Hz), 7.218-7.271 (m, 2H) ppm.

EXAMPLE 51

(2-fluorophenyl)-[(4aS,9bR)-5-methyl-6-trifloromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl]-amine

The title compound was prepared by following the general coupling procedure as a yellow solid (29 mg, 99%) from (4aS,9bR)-8-bromo-5-methyl-6-trifluomethyl-

1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 45, 35 mg, 0.08 mmol) and 2-fluoro-phenylamine (27 mg, 0.24 mmol). ¹H NMR (CDCl₃, 300 MHz) δ 2.006-2.049 (m, 2H), 2.712 (dd, 1H, 9.2, 12.9 Hz), 2.927 (t, 2H, 2.2 Hz), 2.984-3.027 (m, 2H), 3.141 (dd, 1H, 5.8, 12.8 Hz), 3.306-3.337 (m, 1H), 3.499-3.630 (m, 3H), 5.646 (d, 1H, 2.5 Hz), 6.764-6.821 (m, 1H), 6.986-7.101 (m, 4H), 7.211 (d, 1H, 2.2 Hz) ppm.

EXAMPLE 52

10 (3-fluorophenyl)-[(4aS,9bR)-5-methyl-6-trifloromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl)-amine

The title compound was prepared by following the general coupling procedure as a yellow solid (27 mg, 92%) from (4aS,9bR)-8-bromo-5-methyl-6-trifluomethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 45, 35 mg, 0.08 mmol) and 3-fluoro-phenylamine (27 mg, 0.24 mmol). ¹H NMR (CDCl₃, 300 MHz) δ 1.967-2.085 (m, 2H), 2.713 (dd, 1H, 9.1, 12.8 Hz), 2.922-20 2.935 (m, 2H), 2.974-3.032 (m, 2H), 3.152 (dd, 1H, 6.2, 13.2 Hz), 3.305-3.378 (m, 3H), 3.487-3.558 (m, 1H), 5.611 (s, 1H), 6.493-6.617 (m, 3H), 7.055 (d, 1H, 1.8 Hz), 7.117-7.191 (m, 2H) ppm.

(4-fluorophenyl)-[(4aS,9bR)5-methyl-6-trifloromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl]-amine

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The title compound was prepared by following the general coupling procedure as a yellow solid (25 mg, 86%) from (4aS,9bR)-8-bromo-5-methyl-6-trifluomethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 45, 35 mg, 0.08 mmol) and 4-Fluoro-phenylamine (27 mg, 0.24 mmol). ¹H NMR (CDCl₃, 300 MHz) δ 1.874-1.961 (m, 2H), 2.659-2.995 (m, 7H), 3.056 (dd, 1H, 5.9, 12.5 Hz), 3.207 (dd, 1H, 6.2, 14.6 Hz), 3.452-3.504 (m, 1H), 5.386 (s, 1H), 6.801-6.868 (m, 2H), 6.884-6.964 (m, 3H), 7.065 (d, 1H, 2.2 Hz) ppm.

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EXAMPLE 54

(2-ethoxyphenyl)-[(4aS,9bR)-5-methyl-6-trifloromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl)-amine

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The title compound was prepared by following the general coupling procedure as a yellow solid (30 mg, 96%) from (4aS,9bR)-8-bromo-5-methyl-6-trifluomethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester

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(Example 45, 35 mg, 0.08 mmol) and 2-ethoxy-phenylamine (33 mg, 0.24 mmol). 1 H NMR (CDCl₃, 300 MHz) δ 1.460 (t, 3H, 7.0 Hz), 1.891-1.943 (m, 2H), 2.679-2.954 (m, 7H), 3.064 (dd, 1H, 6.3, 12.9 Hz), 3.199-3.226 (m, 1H), 3.483-3.505 (m, 1H), 4.106 (dd, 2H, 6.9, 13.9 Hz), 5.970 (s, 1H), 6.757-6.861 (m, 3H), 6.963 (dd, 1H, 1.8, 8.1 Hz), 7.084 (d, 1H, 2.2 Hz), 7.209 (d, 1H, 2.2 Hz) ppm.

EXAMPLE 55

[(4aS,9bR)-5-methyl-6-trifloromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl]-o-tolyl-amine

The title compound was prepared by following the general coupling procedure as a yellow solid (16 mg, 57%) from (4aS,9bR)-8-bromo-5-methyl-6-trifluomethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 45, 35 mg, 0.08 mmol) and *o*-tolylamine (26 mg, 0.24 mmol). ¹H NMR (CDCl₃, 300 MHz) δ 2.115-2.162 (m, 2H), 2.255 (s, 3H), 2.664 (dd, 1H, 9.9, 12.8 Hz), 2.908 (d, 3H, 1.9 Hz), 3.068-3.223 (m, 3H), 3.361-3.505 (m, 2H), 5.255 (s, 1H), 6.840-6.931 (m, 1H), 6.964-6.986 (m, 2H), 7.055-7.217 (m, 3H) ppm.

(3-fluoro-4-methylphenyl)-[(4aS,9bR)-5-methyl-6-trifloromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl]-amine

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The title compound was prepared by following the general coupling procedure as a yellow solid (21 mg, 69%) from (4aS,9bR)-8-bromo-5-methyl-6-trifluomethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 45, 35 mg, 0.08 mmol) and 3-fluoro-4-methyl-phenylamine (30 mg, 0.24 mmol). ¹H NMR (CDCl₃, 300 MHz) δ 1.886-2.032 (m, 2H), 2.196 (d, 3H, 1.5 Hz), 2.682-3.017 (m, 7H), 3.089 (dd, 1H, 6.2, 12.8 Hz), 3.249 (dd, 1H, 6.2, 14.6 Hz), 3.489-3.543 (m, 1H), 5.480 (s, 1H), 6.525-6.567 (m, 2H), 6.962-7.039 (m, 2H), 7.315 (d, 1H, 2.2 Hz) ppm.

EXAMPLE 57

(3-chloro-4-fluorophenyl)-[(4aS,9bR)-5-methyl-6-trifloromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl]-amine

The title compound was prepared by following the general coupling procedure as a yellow solid (22 mg, 63%) from (4aS,9bR)-8-bromo-5-methyl-6-trifluomethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester

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(Example 45, 35 mg, 0.08 mmol) and 3-chloro-4-fluoro-phenylamine (35 mg, 0.24 mmol). ¹H NMR (CDCl₃, 300 MHz) δ 1.978-1.990 (m, 2H), 2.184-2.221 (m, 3H), 2.663 (dd, 1H, 9.2, 12.4 Hz), 2.887 (d, 2H, 1.8 Hz), 2.952-2.995 (m, 1H), 3.093 (dd, 1H, 5.8, 12.8 Hz), 3.236-3.488 (m, 2H), 5.336 (s, 1H), 6.649-6.757 (m, 2H), 6.847-6.933 (m, 2H), 7.059 (d, 1H, 2.2 Hz) ppm.

EXAMPLE 58

(4-fluoro-3-methylphenyl)-[(4aS,9bR)-5-methyl-6-trifloromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl]-amine

The title compound was prepared by following the general coupling procedure
as a yellow solid (20 mg, 67%) from (4aS,9bR)-8-bromo-5-methyl-6-trifluomethyl1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester
(Example 45, 35 mg, 0.08 mmol) and 4-fluoro-3-methyl-phenylamine (30 mg, 0.24 mmol). ¹H NMR (CDCl₃, 300 MHz) δ 1.845-1.990 (m, 2H), 2.399-3.605 (m, 4H),
2.717 (dd, 1H, 8.4, 12.8 Hz), 2.816-2.984 (m, 5H), 3.056 (dd, 1H, 5.9, 12.8 Hz),
3.176-3.247 (m, 1H), 3.492-3.546 (m, 1H), 5.394 (s, 1H), 6.647-6.699 (m, 1H), 6.822-6.914 (m, 1H), 6.951-7.008 (m, 2H), 7.088 (d, 1H, 2.2 Hz) ppm.

(4-Chloro-3-methylphenyl)-[(4aS,9bR)-5-methyl-6-trifloromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl]-amine

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The title compound was prepared by following the general coupling procedure as a yellow solid (19 mg, 58%) from (4aS,9bR)-8-bromo-5-methyl-6-trifluomethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 45, 35 mg, 0.08 mmol) and 4-chloro-3-methyl-phenylamine (34 mg, 0.24 mmol). ¹H NMR (CDCl₃, 300 MHz) δ 1.886-2.027 (m, 2H), 2.294 (s, 3H), 2.657-2.967 (m, 7H), 3.074 (dd, 1H, 6.3, 12.9 Hz), 3.245 (dd, 1H, 6.6, 15.0 Hz), 3.471-3.524 (m, 1H), 5.417 (s, 1H), 6.635 (dd, 1H, 2.6, 8.4 Hz), 6.712 (d, 1H, 2.5 Hz), 6.976 (d, 1H, 1.8 Hz), 7.110-7.162 (m, 2H) ppm.

EXAMPLE 60

(4aS,9bR)-(2-methoxylphenyl)-(5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl)-amine

The title compound was prepared by following the general coupling procedure as a yellow solid (22 mg, 61%) from (4aS,9bR)-8-bromo-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester

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(Example 45, 41 mg, 0.095 mmol) and 2-methoxyaniline (37 mg, 0.30 mmol). ^{1}H NMR (CDCl₃, 300 MHz) δ (ppm) 1.82-2.00 (m, 2H), 2.58-2.80 (m, 3H), 2.80-3.00 (m, 5H), 3.07 (dd, J = 12.9, 5.9 Hz, 1H), 3.15-3.25 (m, 1H), 3.45-3.55 (m, 1H), 3.91 (s, 3H), 5.97 (s, 1H), 6.75-6.90 (m, 3H), 6.99 (dd, J = 1.6, 7.5 Hz, 1H), 7.09 (d, J = 1.8 Hz, 1H).

EXAMPLE 61

(4aS,9bR)-(2,6-dimethylphenyl)-(5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl)-amine

The title compound was prepared by following the general coupling procedure as a yellow solid (18 mg, 50 %) from (4aS,9bR)-bromo-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 45, 41 mg, 0.095 mmol) and 2,6-dimethylaniline (39 mg, 0.30 mmol). 1 H NMR (CDCl₃, 300 MHz) δ (ppm) 1.87-1.98 (m, 2H), 2.20 (s, 6H), 2.30-2.70 (m, 3H), 2.78-3.20 (m, 6H), 3.33-3.45 (m, 1H), 5.07 (s, 1H), 6.46 (d, J = 2.2 Hz, 1H), 6.59 (d, J = 2.2 Hz, 1H), 7.00-7.18 (m, 3H).

EXAMPLE 62

(4aS,9bR)-(2,4-Difluoro-phenyl)-(5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl)-amine

The title compound was prepared by following the general coupling procedure as a yellow solid (21 mg, 55 %) from (4aS,9bR)-bromo-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 45, 41 mg, 0.095 mmol) and 2,4-difluoroaniline (39 mg, 0.30 mmol). MS (APCI): 384 (base, M+H).

EXAMPLE 63

10 (4aS,9bR)-(2,6-Difluoro-phenyl)-(5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl)-amine

The title compound was prepared by following the general coupling procedure as a yellow solid (12 mg, 31 %) from (4aS,9bR)-bromo-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 45, 41 mg, 0.095 mmol) and 2,6-difluoroaniline (39 mg, 0.30 mmol). MS (APCI): 384 (base, M+H).

EXAMPLE 64

cis-(4a,9b)-8-hydroxy-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid tert-butyl ester

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To a solution of 8-bromo-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 44, 12.6 g, 29 mmol) in anhydrous THF (120 mL) at –78 °C was added *n*-BuLi (2.5 M in hexanes, 17.6 mL, 44 mmol). The reaction was warmed to –60 °C for 45 min. The mixture was cooled to –78 °C and triisopropyl borate (13.5 mL, 58 mmol) was added dropwise. The reaction was stirred at –78 °C for 30 min and then at 0 °C for an additional 20 min. Acetic acid (7.4 mL, 129 mmol) was added followed by the addition of hydrogen peroxide (35 wt % in water, 4.5 mL, 47 mmol). The reaction was stirred at room temperature for 20 min. The mixture was partitioned between 1:1 ether/water (1000 mL). The ether layer was then washed successively with 10% aq sodium thiosulfate (100 mL), water (100 mL), and brine (100 mL), dried over sodium sulfate, and concentrated in vacuo. Purification of the residue by column chromatography (silica gel, 2:1 hexanes/EtOAc) followed by trituration in ether provided the title compound (7.2 g, 67%) as a white solid: MS (ESI): 373 (base, M+H).

EXAMPLE 65

(4aS,9bR)-8-hydroxy-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester

The title compound was prepared by following the above procedure for Example 70 as a white solid from (4aS,9bR)-bromo-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 45). MS (ESI): 373 (base, M+H).

cis-(4a,9b)-5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-ol

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A solution of *cis*-(4a,9b)-8-hydroxy-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 70, 150 mg, 0.4 mmol) in 20% trifluoroacetic acid in dichloromethane (2.5 mL) was stirred for 1h at rt. The reaction mixture was then concentrated *in vacuo*. The residue was dissolved in CHCl3 and washed with 1M NaOH. The organic layer was dried over MgSO4, filtered, and concentrated *in vacuo* to give the title compound as a pale yellow solid (103 mg, 95%) MS (ESI): 273 (base, M+H).

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EXAMPLE 67

(4aS,9bR)-5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-ol

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The title compound was prepared by following the above procedure for Example 72 as a pale yellow solid from (4aS,9bR)-8-hydroxy-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 71). MS (ESI): 273 (base, M+H).

cis-(4a,9b)-8-cyclopropylmethoxy-5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole

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Step A. To a suspension of NaH (60% in mineral oil, 30 mg, 0.75 mmol) in DMF (0.5 mL) at 0 °C was added dropwise a solution of cis-(4a,9b)-8-hydroxy-5-10 methyl-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid tert-butyl ester (Example 70, 162 mg, 0.44 mmol) in DMF (0.5 mL). The reaction mixture was stirred for 30 min at 0 °C then cyclopropylmethyl bromide (60 μL, 0.60 mmol) was added. The ice bath was removed and the reaction mixture was stirred for 1h. The reaction was quenched with the careful addition of crushed ice, 15 then water (2 mL). The mixture was extracted with EtOAc (25 mL) and the organic layer was washed with water (10 mL) and brine (10 mL), dried over anhydrous Na₂SO₄, filtered, and evaporated. Purification of the residue by column chromatography (silica gel, 2:1 hexanes/Et₂O) provided cis-(4a,9b)-8cyclopropylmethoxy-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-20 b]indole-2-carboxylic acid tert-butyl ester (125 mg, 67%) as an oil: ¹H NMR (300 MHz, CDCl₃) δ 6.89–6.82 (m, 2H), 3.89–3.73 (m, 1H), 3.72 (d, J = 6.9 Hz, 2H), 3.61-3.23 (m, 3H), 2.91-2.83 (m, 3H), 1.96-1.68 (m, 2H), 1.44 (s, 9H), 0.68-0.57 (m, 2H), 0.36–0.27 (m, 2H); MS (ESI): 427 (Base, M+H).

Step B. To a stirred solution of *cis*-(4a,9b)-8-cyclopropylmethoxy-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (109 mg, 0.25 mmol) in CH₂Cl₂ (5 mL) was added TFA (0.50 mL). The reaction was stirred at room temperature for 3h and the solvent was evaporated under vacuum. The residue was dissolved in CH₂Cl₂ and the solution was washed with sat. NaHCO₃, dried over Na₂SO₄, filtered, and concentrated in vacuo. Purification of the

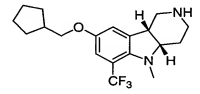
residue by CombiFlash chromatography [silica gel, 0–20% (CHCl3/MeOH/NH4OH)/EtOAc] provided the title compound (48 mg, 58%) as a light yellow oil: 1 H NMR (300 MHz, CD3OD) δ 6.94 (d, J = 2.3 Hz, 1H), 6.83 (d, J = 2.5 Hz, 1H), 3.75 (d, J = 6.8 Hz, 1H), 3.43–3.34 (m, 1H), 3.18–3.07 (m, 1H), 2.95 (dd J = 6.2, 12.9 Hz, 1H), 2.89–2.71 (m, 5H), 2.48 (dd, J = 9.2, 12.9 Hz, 1H), 1.98–1.75 (m, 2H), 1.28–1.12 (m, 1H), 0.63–0.54 (m, 2H), 0.36–0.27 (m, 2H); MS (ESI): 327 (Base M+H).

EXAMPLE 69

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cis-(4a,9b)-8-cyclopentylmethoxy-5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole



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Following the procedure described for Example 74 Step A, and B, the title compound was prepared as a yellow oil in 57% yield: 1 H NMR (300 MHz, CD₃OD) δ 6.93 (d, J = 2.3 Hz, 1H), 6.82 (d, J = 2.5 Hz, 1H), 3.78 (d, J = 6.9 Hz, 2H), 3.42–3.37 (m, 1H), 3.19–3.08 (m, 1H), 2.96 (dd, J = 12.9, 6.1 Hz, 1H), 2.89–2.74 (m, 5H), 2.48 (dd, J = 12.9, 9.1 Hz, 1H), 2.39–2,24 (m, 1H), 1.99–1.74 (m, 4H), 1.74–1.59 (m, 4H), 1.47–1.28 (m, 2H); ESI MS m/z 355 [$C_{19}H_{25}F_{3}N_{2}O + H$] $^{+}$.

cis-(4a,9b)-5-methyl-8-(3-methyl-butoxy)-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole

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Following the procedure described for Example 74 Step A, cis-(4a,9b)-5-methyl-8-(3-methyl-butoxy)-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid tert-butyl ester was prepared using Example 70 (124 mg, 0.33 mmol), 3-methylbromobutane (60 μ L, 0.48 mmol), and NaH (60% in mineral oil, 27 mg, 0.68 mmol) in 76% yield (111 mg) as an oil: 1 H NMR (300 MHz, CD3OD) δ 6.95 (d, J = 2.4 Hz, 1H), 6.80 (d, J = 2.4 Hz, 1H), 3.96–3.91 (m, 2H), 3.79–3.12 (m, 6H), 2.89–2.83 (m, 3H), 2.04–1.73 (m, 3H), 1.68–1.57 (m, 2H), 1.36 (s, 9H), 0.95 (d, J = 6.8 Hz, 6H); ESI MS m/z 443 [C₂₃H₃₃F₃N₂O₃ + H]⁺.

The title compound was prepared by following the deprotection procedure of Example 74 Step B as a as a yellow oil: 1 H NMR (300 MHz, CD₃OD) δ 6.94 (d, J = 2.3 Hz, 1H), 6.83 (d, J = 2.5 Hz, 1H), 3.94 (t, J = 6.5 Hz, 1H), 3.44–3.35 (m, 1H), 3.19–3.09 (m, 1H), 2.95 (dd, J = 12.9, 6.1 Hz, 1H), 2.86–2.73 (m, 5H), 2.48 (dd, J = 12.9, 9.1 Hz, 1H), 1.98–1.77 (m, 3H), 1.62 (q, J = 6.6 Hz, 2H), 0.96 (d, J = 6.7 Hz, 6H); ESI MS m/z 343 $[C_{17}H_{25}F_{3}N_{2}O + H]^{+}$.

cis-(4a,9b)-5-methyl-8-propoxy-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole

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Following the procedure described for Example 74 Step A, cis-(4a,9b)-5-methyl-8-propoxy-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid tert-butyl ester was prepared using Example 70 (130 mg, 0.35 mmol), bromopropane (40 μ L, 0.44 mmol), and NaH (60% in mineral oil, 30 mg, 0.75 mmol) in 83% yield (120 mg) as an oil: 1 H NMR (300 MHz, CD₃OD) δ 6.95 (d, J = 2.4 Hz, 1H), 6.80 (d, J = 2.4 Hz, 1H), 3.92–3.80 (m, 2H), 3.75–3.14 (m, 6H), 2.91–2.85 (m, 3H), 2.02–1.68 (m, 4H), 1.37 (s, 9H), 1.03 (t, J = 7.4 Hz, 3H); ESI MS m/z 415 [C₂₁H₂₉F₃N₂O₃ + H]⁺.

The title compound was prepared by following the deprotection procedure of Example 74 Step B as a as a light yellow oil in 52% yield: 1 H NMR (300 MHz, CD₃OD) δ 6.94 (d, J = 2.4 Hz, 1H), 6.84 (d, J = 2.5 Hz, 1H), 3.89 (t, J = 6.4 Hz, 1H), 3.47–3.27 (m, 1H), 3.21–3.09 (m, 1H), 2.96 (dd, J = 12.9, 6.1 Hz, 1H), 2.86–2.70 (m, 5H), 2.48 (dd, J = 12.9, 9.2 Hz, 1H), 2.00–1.68 (m, 4H), 1.02 (t, J = 7.4 Hz, 3H); ESI MS m/z 315 [C₁₆H₂₁F₃N₂O + H]⁺.

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cis-(4a,9b)-8-butoxy-5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole

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Following the procedure described for Example 74 Step A, cis-(4a,9b)-5-methyl-8-butoxy-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid tert-butyl ester was prepared using Example 70 (122 mg, 0.33 mmol), bromobutane (50 μ L, 0.46 mmol), and NaH (60% in mineral oil, 37 mg, 0.92 mmol) in 84% yield (119 mg) as an oil: 1 H NMR (300 MHz, CD₃OD) δ 6.95 (d, J = 2.4 Hz, 1H), 6.80 (d, J = 2.4 Hz, 1H), 3.98–3.87 (m, 2H), 3.76–3.16 (m, 6H), 2.91–2.85 (m, 3H), 2.05–1.65 (m, 4H), 1.56–1.42 (m, 2H), 1.36 (s, 9H), 0.98 (t, J = 7.4 Hz, 3H); ESI MS m/z 429 [$C_{22}H_{31}F_{3}N_{2}O_{3} + H$]⁺.

The title compound was prepared by following the deprotection procedure of Example 74 Step B as a as a light yellow oil in 44% yield: 1 H NMR (300 MHz, CD₃OD) δ 6.94 (d, J = 2.4 Hz, 1H), 6.83 (d, J = 2.5 Hz, 1H), 3.91 (t, J = 6.3 Hz, 1H), 3.44–3.36 (m, 1H), 3.19–3.10 (m, 1H), 2.96 (dd, J = 12.9, 6.2 Hz, 1H), 2.89–2.74 (m, 5H), 2.49 (dd, J = 12.8, 9.2 Hz, 1H), 1.99–1.80 (m, 2H), 1.58–1.42 (m, 2H), 0.98 (t, J = 7.3 Hz, 3H); ESI MS m/z 329 [C₁₇H₂₃F₃N₂O + H]⁺.

cis-(4a,9b)-8-(3,3-dimethyl-butoxy)-5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole

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$$O$$
 CF_3
 N
 H
 NH

Following the procedure described for Example 74 Step A, and B, the title compound was prepared as a light yellow oil in 53% yield: 1 H NMR (300 MHz, CD₃OD) δ 6.93 (d, J = 2.4 Hz, 1H), 6.82 (d, J = 2.6 Hz, 1H), 3.98 (t, J = 6.9 Hz, 1H), 3.43–3.37 (m, 1H), 3.19–3.09 (m, 1H), 2.96 (dd, J = 12.8, 6.1 Hz, 1H), 2.88–2.73 (m, 5H), 2.49 (dd, J = 12.8, 9.2 Hz, 1H), 1.98–1.77 (m, 2H), 1.68 (t, J = 7.0 Hz, 2H), 0.99 (s, 9H); ESI MS m/z 357 [C₁₉H₂₇F₃N₂O + H]⁺.

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EXAMPLE 74

cis-(4a,9b)-8-cyclobutylmethoxy-5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole

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Following the procedure described for Example 74 Step A, cis-(4a,9b)-5-methyl-8-cyclobutylmethoxy-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid tert-butyl ester was prepared using Example 70 (122 mg, 0.33 mmol), cyclobutylmethyl bromide (44 μ L, 0.40 mmol), and NaH (60% in mineral oil, 16 mg, 0.40 mmol) in 36% yield (52 mg) as an oil: 1 H NMR (300 MHz, CD3OD) δ 6.96 (d, J = 2.5 Hz, 1H), 6.80 (d, J = 2.5 Hz, 1H), 3.92–3.81 (m, 2H),

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3.78-3.32 (m 7H), 3.28-3.13 (m 1H), 2.91-2.84 (m, 3H), 2.79-2.65 (m, 1H), 2.20-1.76 (m, 6H), 1.37 (s, 9H); ESI MS m/z 441 [C₂₃H₃₁F₃N₂O₃ + H]⁺.

The title compound was prepared by following the deprotection procedure of Example 74 Step B as a as a light yellow oil in 33% yield: ¹H NMR (300 MHz, CD₃OD δ 6.94 (d, J = 2.4 Hz, 1H), 6.83 (d, J = 2.5 Hz, 1H), 3.87 (d, J = 6.4 Hz, 1H), 3.43–3.35 (m, 1H), 3.29–3.19 (m, 1H), 2.96 (dd, J = 12.9, 6.2 Hz, 1H), 2.89–2.64 (m, 6H), 2.49 (dd, J = 12.9, 9.2 Hz, 1H), 2.18–2.04 (m, 2H), 2.04–1.78 (m, 6H); ESI MS m/z 341 [C₁₈H₂₃F₃N₂O + H]⁺.

General method for preparation of 5-methyl-8-(pyridin-yl-methoxy)-6trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole

8-Hydroxy-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (1.0 eq.), bromomethylpyridine (2.0 eq.), and K₂CO₃ (4.0 eq.) were dissolved in anhydrous DMF(0.3 M). The mixture was stirred under Ar at rt for 24 h. Same work up and deprotection procedure were followed as described in Example 74 Method A and B to obtain the title compound.

EXAMPLE 75

20 (4aS,9bR)-5-methyl-8-(pyridin-2-ylmethoxy)-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole

The title compound was prepared by following the general method as a yellow solid (60 mg, 61%) from (4aS,9bR)-8-hydroxy-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 71, 100 mg, 0.27mmol), 2-bromomethyl-pyridine (141 mg, 0.56 mmol) and K2CO3 (156 mg, 1.12 mmol). MS (ESI): 364 (base, M+H).

EXAMPLE 76

(4aS,9bR)-5-methyl-8-(pyridin-3-ylmethoxy)-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole

The title compound was prepared by following the general method as a white solid (48 mg, 49%) from (4aS,9bR)-8-hydroxy-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 71, 100 mg, 0.27mmol), 3-bromomethyl-pyridine (141 mg, 0.56 mmol) and K2CO3 (156 mg, 1.12 mmol). MS (ESI): 364 (base, M+H).

15 EXAMPLE 77

(4aS,9bR)-5-methyl-8-(pyridin-4-ylmethoxy)-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole

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The title compound was prepared by following the general Method C as a light yellow solid (56 mg, 57%) from (4aS,9bR)-8-hydroxy-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 71, 100 mg, 0.27mmol), 4-bromomethyl-pyridine (141 mg, 0.56 mmol) and K2CO3 (156 mg, 1.12 mmol). MS (ESI): 364 (base, M+H).

EXAMPLE 78

cis-(4a,9b)-5-methyl-8-o-tolyloxy-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole

To a solution of *cis*-(4a,9b)-8-hydroxy-5-methyl-6-trifluoromethyl1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester
(Example 70, 50 mg, 0.13 mmol) in triethylamine (38 μL, 0.27 mmol) and CH₂Cl₂
was added 2-methylphenyl boronic acid (37 mg, 0.27 mmol) and Cu(OAc)₂ (36 mg, 0.2 mg). The reaction mixture was stirred open to the air at rt for 15 h. The reaction mixture was filtered and concentrated in vacuo. The residue was chromatographed
(silica gel, Hex/EtoAc 0-30%) to give 5-methyl-8-o-tolyloxy-6-trifluoromethyl1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid tert-butyl ester.
Deprotection of Boc group was followed as described in Example 74 to obtain the title compound as a yellow oil (13 mg, 28%). MS (ESI): 364 (Base M+H).

20 **EXAMPLE 79**

cis-(4a,9b)-8-(2,5-dimethyl-phenoxy)-5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole

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Following the procedure described for Example 84 Step A, the title compound was prepared using *cis*-(4a,9b)-8-hydroxy-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 70, 50 mg, 0.13 mmol), triethylamine (38 μL, 0.27 mmol), 2,5-dimethylphenyl boronic acid (41 mg, 0.27 mmol) and Cu(OAc)₂ (36 mg, 0.2 mg) as a yellow oil (10 mg, 20%). MS (ESI): 377 (Base M+H).

EXAMPLE 80

10 cis-(4a,9b)-2-(5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yloxy)-benzonitrile

A mixture of *cis*-(4a,9b)-8-hydroxy-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 70, 50 mg, 0.13 mmol), 2-fluoro-benzylcarbonitrile (22 μL, 0.2 mmol), K₂CO₃ (29 mg, 0.21 mmol) in DMF (1 ml) was irradiated in microwave for 900 sec. at 160 °C. The reaction mixture was partitioned between H₂O and EtOAc. The organic layer was washed with H₂O, dried over MgSO₄, filtered and concentrated in vacuo. The residue was chromatographed (silica gel, Hex/EtoAc 0-50%) to give 8-(2-cyano-phenoxy)-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester. Deprotection of Boc group was followed as described in Example 74 to obtain the title compound as a yellow oil (21mg, 43%). MS (ESI): 374 (Base M+H).

cis-(4a,9b)-4-(5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yloxy)-benzonitrile

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Following the procedure described for Example 84 Step A, the title compound was prepared using *cis*-(4a,9b)-8-hydroxy-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 70, 50 mg, 0.13 mmol), triethylamine (38 μL, 0.27 mmol), 4-cyano-phenyl boronic acid (39 mg, 0.27 mmol) and CuF₆(MeCN)₄ (36 mg, 0.2 mg) as a yellow oil (12 mg, 25%). MS (ESI): 374 (Base M+H).

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EXAMPLE 82

cis-(4a,9b)-8-(2-methoxy-phenoxy)-5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole

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Following the procedure described for Example 84 Step A, the title compound was prepared using cis-(4a,9b)-8-hydroxy-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid tert-butyl ester (Example 70, 50 mg, 0.13 mmol), triethylamine (38 μ L, 0.27 mmol), 2-methoxy-phenyl boronic acid (41 mg, 0.27 mmol) and CuF₆(MeCN)₄ (36 mg, 0.2 mg) as a yellow oil (11 mg, 22%). MS (ESI): 388 (Base M+H).

EXAMPLE 83

cis-(4a,9b)-8-amino-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydropyrido[4,3-b]indole-2-carboxylic acid tert-butyl ester

A solution of cis-(4a,9b)-8-bromo-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-10 hexahydro-pyrido[4,3-b]indole-2-carboxylic acid tert-butyl ester. (Example 44) (434 mg, 1.0 mmol), benzophenone imine (218 mg, 1.2 mmol), (S)-(-)-2,2'bis(diphenylphosphino)-1,1'-binaphthyl (BINAP) (46.7 mg, 0.075 mmol), Pd2dba3 (46 mg, 0.05 mmol) and sodium tert-butoxide (135 mg, 1.4 mmol) in degassed toluene (5 mL) was heated at reflux under Argon atmosphere for 15 h. The solution 15 was cooled, and H2O was added to the reaction mixture. The mixture extracted with CH₂Cl₂. The organic layer was dried over MgSO₄, filtered and concentrated in vacuo. The residue was taken up in methanol (15 mL) and then there was added NaOAC (197 mg) and hydroxylamine hydrochloride (127 mg) and the mixture was stirred at rt for 3 h. To the reaction mixture was diluted with CH2Cl2 then washed 20 with H₂O. The organic solution was dried over MgSO₄, filtered and concentrated in vacuo. The residue was purified by column chromatography (EtOAc/hexane 2/8) to afford the title compound (269 mg, 73%) as a yellow solid. MS (APCI): 372 (base, M+H).

(4aS,9bR)-8-amino-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydropyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester

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The title compound was prepared by following the procedure described for Example 89 using (4aS,9bR)-8-bromo-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 45). MS (APCI): 372 (base, M+H).

EXAMPLE 85

cis-(4a,9b)-5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-ylamine

1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 89, 100 mg, 0.27 mmol) in CH₂Cl₂ (5 mL)was added TFA (1 mL) at rt. The reaction mixture was stirredd for 1 h at rt then concentrated *in vacuo*. The

organic solution was dried over MgSO₄, filtered and concentrated *in vacuo*. to yield the title compound (70 mg, 95%) as a yellow solid: MS (ESI): 272 (base, M+H).

residue was dissolved in CHCl3 (20 mL) and washed with 1M NaOH (5 mL). The

To a solution of cis-(4a,9b)-8-amino-5-methyl-6-trifluoromethyl-

(4aS,9bR)-5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-ylamine

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$$H_2N$$
 N
 N
 H

The title compound was prepared by following the procedure described for Example 91 using (4aS,9bR)-8-amino-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 90). MS (APCI): 272 (base, M+H).

General coupling method for preparation of (5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl)-pyridin-3-yl-amine

Method A. 8-bromo-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydropyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 44 or45, 1.0 eq.), aminopyridine (3.0 eq.), and NaOt-Bu (3.0 eq.) were dissolved in anhydrous toluene (0.17 M). The mixture was degassed with argon for 30 min. Pd2(dba)3 (0.06 eq.) and 2,2'-bis(diphenylphosphino)-1,1'-binaphthyl (0.18 eq.) were added; the reaction was heated at 80 °C for 16 h. The reaction was cooled to room temperature, diluted with EtOAc, filtered through a bilayer pad of diatomaceous earth and silica gel, and concentrated. Purification of the residue by flash column chromatography (silica gel, 2–30% EtOAc/hexanes) provided 5-methyl-8-(pyridinylamino)-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid tert-butyl ester derivatives. The intermediate was dissolved in CH2Cl2/TFA (5/1) at rt stirred for 1 h. Upon concentration in vacuo, the residue was partitioned between CH2Cl2/1 M NaOH. The aqueous phase was extracted with CH2Cl2. The combined organic phases were dried over MgSO4, filtered, and concentrated *in vacuo*. The residue was purified by flash column chromatography to give the title compound.

Method B. 8-amino-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydropyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 91 or 92, 1.0 eq.), bromopyridine (1.0 eq.), and CsCO₃ (2.0 eq.) were dissolved in anhydrous toluene (0.1 M). The mixture was degassed with argon for 30 min. Pd₂(dba)₃ (0.01 eq.) and 2,2'-bis(diphenylphosphino)-1,1'-binaphthyl (0.13 eq.) were added; the reaction was heated at 80 °C for 16 h. Same work up and deprotection procedure were followed as described in Method A to obtain the title compound.

EXAMPLE 87

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cis-(4a,9b)-(5-Methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl)-pyridin-3-yl-amine

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The title compound was prepared by following the general Method A as a yellow solid (24 mg, 14%) from *cis*-(4a,9b)-8-bromo-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 44, 217 mg, 0.5 mmol), 3-amino-pyridine (141 mg, 1.5 mmol) and NaO*t*-Bu (144 mg, 1.5 mmol). MS (ESI): 349 (base, M+H).

EXAMPLE 88

cis-(4a,9b)-(2-chloro-pyridin-3-yl)-(5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl)-amine

The title compound was prepared by following the general Method A as a yellow solid (68 mg, 36%) from *cis*-(4a,9b)-8-bromo-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 44, 217 mg, 0.5 mmol), 3-amino-2-chloro-pyridine (192 mg, 1.5 mmol) and NaO*t*-Bu (144 mg, 1.5 mmol). MS (ESI): 383 (base, M+H).

EXAMPLE 89

10 cis-(4a,9b)-3-(5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-ylamino)-pyridine-2-carbonitrile

The title compound was prepared by following the general Method B as a yellow solid (67 mg, 60%) from *cis*-(4a,9b)-8-amino-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 89, 111 mg, 0.3 mmol), 3-bromo-2-cyano-pyridine (55 mg, 0.3 mmol) and CsCO₃ (196 mg, 0.6 mmol). MS (ESI): 374 (base, M+H).

EXAMPLE 90

(4aS,9bR)-3-(5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-ylamino)-pyridine-2-carbonitrile

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The title compound was prepared by following the general Method B as a yellow solid from (4aS,9bR)-8-amino-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 90). MS (ESI): 374 (base, M+H).

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EXAMPLE 91

cis-(4a,9b)-(6-methoxy-pyridin-3-yl)-(5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl)-amine

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The title compound was prepared by following the general Method A as a yellow solid (52 mg, 28%) from *cis*-(4a,9b)-8-bromo-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 44, 217 mg, 0.5 mmol), 5-amino-2-methoxy-pyridine (186 mg, 1.5 mmol) and NaO*t*-Bu (144 mg, 1.5 mmol). MS (ESI): 379 (base, M+H).

EXAMPLE 92

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(4aS,9bR)-(6-methoxy-pyridin-3-yl)-(5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl)-amine

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The title compound was prepared by following the general Method A as a yellow solid (85 mg, 45%) from (4aS,9bR)-8-bromo-5-methyl-6-trifluoromethyl-

1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 45, 217 mg, 0.5 mmol), 5-amino-2-methoxy-pyridine (186 mg, 1.5 mmol) and NaO*t*-Bu (144 mg, 1.5 mmol). MS (ESI): 379 (base, M+H).

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EXAMPLE 93

cis-(4a,9b)-(6-fluoro-pyridin-3-yl)-(5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl)-amine

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The title compound was prepared by following the general Method A as a yellow solid (14 mg, 8%) from *cis*-(4a,9b)-8-bromo-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 44, 217 mg, 0.5 mmol), 5-amino-2-fluoro-pyridine (168 mg, 1.5 mmol) and NaO*t*-Bu (144 mg, 1.5 mmol). MS (ESI): 367 (base, M+H).

EXAMPLE 94

20 (4aS,9bR)-(6-fluoro-pyridin-3-yl)-(5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl)-amine

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The title compound was prepared by following the general Method A as a yellow solid from (4aS,9bR)-8-bromo-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-

hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 45). MS (ESI): 367 (base, M+H).

EXAMPLE 95

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cis-(4a,9b)-5-(5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-ylamino)-pyridine-2-carbonitrile

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The title compound was prepared by following the general Method A as a yellow solid (10 mg, 6%) from *cis*-(4a,9b)-8-bromo-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 44, 217 mg, 0.5 mmol), 5-amino-2-cyano-pyridine (179 mg, 1.5 mmol) and NaO*t*-Bu (144 mg, 1.5 mmol). MS (ESI): 374 (base, M+H).

EXAMPLE 96

cis-(4a,9b)- 5-(5-Methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-ylamino)-nicotinonitrile

The title compound was prepared by following the general Method B as a yellow solid (33 mg, 18%) from *cis*-(4a,9b)-8-amino-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester

(Example 89, 186 mg, 0.5 mmol), 5-bromo-3-cyano-pyridine (92 mg, 0.5 mmol) and CsCO₃ (326 mg, 1.0 mmol). MS (ESI): 374 (base, M+H).

EXAMPLE 97

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cis-(4a,9b)-5-(5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-ylamino)-nicotinic acid methyl ester

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The title compound was prepared by following the general Method B as a yellow solid (29 mg, 14%) from *cis*-(4a,9b)-8-amino-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 89, 186 mg, 0.5 mmol), 5-bromo-nicotinic acid methyl ester (108 mg, 0.5 mmol) and CsCO₃ (326 mg, 1.0 mmol). MS (ESI): 407 (base, M+H).

EXAMPLE 98

cis-(4a,9b)-5-(5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-ylamino)-nicotinic acid ethyl ester

The title compound was prepared by following the general Method B as a yellow oil (52 mg, 25%) from *cis*-(4a,9b)-8-amino-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester

(Example 89, 186 mg, 0.5 mmol), 5-Bromo-nicotinic acid ethyl ester (115 mg, 0.5 mmol) and CsCO₃ (326 mg, 1.0 mmol). MS (ESI): 421 (base, M+H).

EXAMPLE 99

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(4aS,9bR)-(2-methoxy-pyridin-3-yl)-(5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl)-amine

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The title compound was prepared by following the general Method A as a yellow solid (25 mg, 13%) from (4aS,9bR)-8-bromo-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 45, 217 mg, 0.5 mmol), 3-amino-2-methoxy-pyridine (186 mg, 1.5 mmol) and NaOt-Bu (144 mg, 1.5 mmol). MS (ESI): 379 (base, M+H).

EXAMPLE 100

cis-(4a,9b)- (6-Chloro-pyridin-3-yl)-(5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl)-amine

The title compound was prepared by following the general Method A as a yellow solid (48 mg, 25%) from *cis*-(4a,9b)-8-bromo-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester

(Example 44, 217 mg, 0.5 mmol), 5-amino-2-chloro-pyridine (193 mg, 1.5 mmol) and NaOt-Bu (144 mg, 1.5 mmol). MS (ESI): 383 (base, M+H).

EXAMPLE 101

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(4aS,9bR)- (6-Chloro-pyridin-3-yl)-(5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl)-amine

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The title compound was prepared by following the general Method A as a yellow solid (56 mg, 29%) from (4aS,9bR)-8-bromo-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 45, 217 mg, 0.5 mmol), 5-amino-2-chloro-pyridine (193 mg, 1.5 mmol) and NaO*t*-Bu (144 mg, 1.5 mmol). MS (ESI): 383 (base, M+H).

EXAMPLE 102

cis-(4a,9b)-(2-ethoxy-pyridin-3-yl)-(5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl)-amine

The title compound was prepared by following the general Method B as a yellow solid (12 mg, 7%) from *cis*-(4a,9b)-8-amino-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester

(Example 89, 160 mg, 0.43 mmol), 3-bromo-2-ethoxypyridine (101 mg, 0.5 mmol) and CsCO₃ (326 mg, 1.0 mmol). MS (ESI): 393 (base, M+H).

EXAMPLE 103

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cis-(4a,9b)-(2,6-dimethoxy-pyridin-3-yl)-(5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl)-amine

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The title compound was prepared by following the general Method A as a yellow solid (22 mg, 11%) from *cis*-(4a,9b)-8-bromo-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 44, 217 mg, 0.5 mmol), 3-amino-2,6-dimethoxy-pyridine hydrochloride (286 mg, 1.5 mmol) and NaOt-Bu (144 mg, 1.5 mmol). MS (ESI): 409 (base, M+H).

EXAMPLE 104

(4aS,9bR)-(2,6-dimethoxy-pyridin-3-yl)-(5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl)-amine

The title compound was prepared by following the general Method A as a yellow solid (4aS,9bR)-8-bromo-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 45). MS (ESI): 409 (base, M+H).

EXAMPLE 105

cis-(4a,9b)-(2,6-dichloro-pyridin-3-yl)-(5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl)-amine

The title compound was prepared by following the general Method A as a yellow solid (21 mg, 10%) from *cis*-(4a,9b)-8-bromo-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 44, 217 mg, 0.5 mmol), 3-amino-2,6-dichloropyridine (244 mg, 1.5 mmol) and NaOt-Bu (144 mg, 1.5 mmol). MS (ESI): 417 (base, M+H).

15 **EXAMPLE 106**

cis-(4a,9b)-4-methyl-3-(5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-ylamino)-pyridine-2-carbonitrile

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The title compound was prepared by following the general Method B as a yellow solid (20 mg, 10%) from *cis*-(4a,9b)-8-amino-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 89, 186 mg, 0.5 mmol), 3-bromo-4-methyl-pyridine-2-carbonitrile (99 mg, 0.5 mmol) and CsCO₃ (326 mg, 1.0 mmol). MS (ESI): 388 (base, M+H).

cis-(4a,9b)-(6-fluoro-5-methyl-pyridin-3-yl)-(5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl)-amine

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The title compound was prepared by following the general Method A as a yellow solid (18 mg, 10%) from *cis*-(4a,9b)-8-bromo-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 44, 217 mg, 0.5 mmol), 5-amino-2-fluoro-3-methylpyridine (189 mg, 1.5 mmol) and NaOt-Bu (144 mg, 1.5 mmol). MS (ESI): 381 (base, M+H).

EXAMPLE 108

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(4aS,9bR)-(6-fluoro-5-methyl-pyridin-3-yl)-(5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl)-amine

20

The title compound was prepared by following the general Method A as a yellow solid from (4aS,9bR)-8-bromo-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 45). MS (ESI): 381 (base, M+H).

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cis-(4a,9b)-(2,5-dichloro-pyridin-3-yl)-(5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl)-amine

5

The title compound was prepared by following the general Method A as a yellow solid (22 mg, 11%) from *cis*-(4a,9b)-8-bromo-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 44, 217 mg, 0.5 mmol), 3-amino-2,5-dichloropyridine (245 mg, 1.5 mmol) and NaOt-Bu (144 mg, 1.5 mmol). MS (ESI): 417 (base, M+H).

EXAMPLE 110

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cis-(4a,9b)-5-methoxy-3-(5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-ylamino)-pyridine-2-carbonitrile

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The title compound was prepared by following the general Method B as a yellow solid (142 mg, 70%) from *cis*-(4a,9b)-8-amino-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 89, 186 mg, 0.5 mmol), 3-bromo-2-cyano-5-methoxy-pyridine (107 mg, 0.5 mmol) and CsCO₃ (326 mg, 1.0 mmol). MS (ESI): 404 (base, M+H).

cis-(4a,9b)-(2-methoxy-6-methyl-pyridin-3-yl)-(5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl)-amine

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The title compound was prepared by following the general Method B as a yellow solid (42 mg, 25%) from *cis*-(4a,9b)-8-amino-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 89, 160 mg, 0.43 mmol), 3-bromo-2-methoxy-6-methylpyridine (101 mg, 0.5 mmol) and CsCO₃ (326 mg, 1.0 mmol). MS (ESI): 393 (base, M+H).

EXAMPLE 112

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cis-(4a,9b)-(6-chloro-2-methyl-pyridin-3-yl)-(5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl)-amine

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25

The title compound was prepared by following the general Method B as a yellow solid (47 mg, 28%) from *cis*-(4a,9b)-8-amino-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 89, 160 mg, 0.43 mmol), 3-bromo-6-chloro-2-methylpyridine (101 mg, 0.5 mmol) and CsCO₃ (326 mg, 1.0 mmol) as a major product. MS (ESI): 397 (base, M+H).

cis-(4a,9b)-6-methyl-3-(5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-ylamino)-pyridine-2-carbonitrile

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The title compound was prepared by following the general Method B as a yellow solid (109 mg, 66%) from *cis*-(4a,9b)-8-amino-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 89, 160 mg, 0.43 mmol), 3-bromo-2-cyano-6-methylpyridine (99 mg, 0.5 mmol) and CsCO₃ (326 mg, 1.0 mmol) as a major product. MS (ESI): 388 (base, M+H).

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EXAMPLE 114

cis-(4a,9b)-(2,6-dimethyl-pyridin-3-yl)-(5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl)-amine

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The title compound was prepared by following the general Method B as a yellow solid (28 mg, 17%) from *cis*-(4a,9b)-8-amino-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 89, 160 mg, 0.43 mmol), 3-bromo-2,6-dimethylpyridine (93 mg, 0.5 mmol) and CsCO₃ (326 mg, 1.0 mmol) as a major product. MS (ESI): 377 (base, M+H).

cis-(4a,9b)-(2-isopropoxy-6-methyl-pyridin-3-yl)-(5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl)-amine

5

The title compound was prepared by following the general Method B as a yellow solid (58 mg, 32%) from *cis*-(4a,9b)-8-amino-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 89, 160 mg, 0.43 mmol), 3-bromo-2-isopropoxy-6-methylpyridine (115 mg, 0.5 mmol) and CsCO3 (326 mg, 1.0 mmol). MS (ESI): 421 (base, M+H).

EXAMPLE 116

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cis-(4a,9b)-(2-ethoxy-6-methyl-pyridin-3-yl)-(5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl)-amine

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The title compound was prepared by following the general Method B as a yellow solid (10 mg, 6%) from *cis*-(4a,9b)-8-amino-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 89, 160 mg, 0.43 mmol), 3-bromo-2-ethoxy-6-methylpyridine (108 mg, 0.5 mmol) and CsCO₃ (326 mg, 1.0 mmol). MS (ESI): 407 (base, M+H).

cis-(4a,9b)-(2-methoxy-4-methyl-pyridin-3-yl)-(5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl)-amine

5

The title compound was prepared by following the general Method B as a yellow solid (15 mg, 9%) from *cis*-(4a,9b)-8-amino-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 89, 160 mg, 0.43 mmol), 3-bromo-2-methoxy-4-methylpyridine (101 mg, 0.5 mmol) and CsCO₃ (326 mg, 1.0 mmol). MS (ESI): 393 (base, M+H).

EXAMPLE 118

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cis-(4a,9b)-isoquinolin-4-yl-(5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl)-amine

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The title compound was prepared by following the general Method B as a yellow solid (51 mg, 43%) from *cis*-(4a,9b)-8-amino-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 89, 111 mg, 0.3 mmol), 4-bromo-isoquinoline (62 mg, 0.3 mmol) and CsCO₃ (196 mg, 0.6 mmol). MS (ESI): 399 (base, M+H).

cis-(4a,9b)-(5-methyl-pyridin-2-yl)-(5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl)-amine

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The title compound was prepared by following the general Method A as a yellow solid (10 mg, 6%) from *cis*-(4a,9b)-8-bromo-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 44, 217 mg, 0.5 mmol), 2-amino-5-methylpyridine (162 mg, 1.5 mmol) and NaOt-Bu (144 mg, 1.5 mmol). MS (ESI): 363 (base, M+H).

EXAMPLE 120

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cis-(4a,9b)-(5-bromo-6-methyl-pyridin-2-yl)-(5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl)-amine

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The title compound was prepared by following the general Method B as a yellow solid (31 mg, 16%) from *cis*-(4a,9b)-8-amino-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 89, 160 mg, 0.43 mmol), 3-bromo-6-chloro-2-methylpyridine (101 mg, 0.5 mmol) and CsCO₃ (326 mg, 1.0 mmol) as a minor product. MS (ESI): 441 (base, M+H).

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General method for preparation of 8-alkyl-5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole

Step A. To a degassed solution of 8-bromo-5-methyl-6-trifluomethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 44 or 45, 1.0 mole equivalent) in DMF (0.067 M) was added alkyl zinc bromide (0.5 M in THF, 2.5 mole equivalent) and Pd(PPh3)4 (0.06 mole equivalent). The reaction mixture was heated at 90°C for 2-5 h, then cooled to rt. The reaction mixture was then quenched with water and extracted with ethyl acetate. The organic layer was dried over MgSO4, filtered and concentrated *in vacuo*. The crude product was chromatographed on a silica gel column by elution with Hexane/Ethyl Acetate to give 8-alkyl-5-methyl-6-trifluoromethyl-1-2,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester in 20-80% yield.

Step B. A solution of 8-alkyl-5-methyl-6-trifluoromethyl-1-2,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid tert-butyl ester in 20% trifluoroacetic acid in CH₂Cl₂ was stirred for 1h at rt. The reaction mixture was then concentrated *in vacuo* and then neutralized with base and extracted with chloroform. The organic layer was concentrated *in vacuo* to give 8-alkyl-5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole (90-100% yield).

20 **EXAMPLE 121**

cis-(4a,9b)-8-(2-ethyl-butyl)-5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole

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The title compound was prepared by following the general coupling procedure as a colorless oil from *cis*-(4a,9b)-8-bromo-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 44) and 2-ethylbutyl zinc bromide. MS (ES+): 341 (base, M+H).

cis-(4a,9b)-8-benzyl-5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole

The title compound was prepared by following the general coupling procedure as a colorless oil from *cis*-(4a,9b)-8-bromo-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 44) and benzyl zinc bromide. MS (ES+): 347 (base, M+H).

EXAMPLE 123

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(4aS,9bR)-8-benzyl-5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole

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The title compound was prepared by following the general coupling procedure as a colorless oil from (4aS,9bR)-8-bromo-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 45) and benzyl zinc bromide. MS (ES+): 347 (base, M+H).

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cis-(4a,9b)-8-cyclohexyl-5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole

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The title compound was prepared by following the general coupling procedure as a colorless oil from *cis*-(4a,9b)-8-bromo-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-10 hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 44) and cyclohexyl zinc bromide. MS (ES+): 339 (base, M+H).

EXAMPLE 125

cis-(4a,9b)-2-(5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-ylmethyl)-benzonitrile

The title compound was prepared by following the general coupling procedure as a colorless oil from *cis*-(4a,9b)-8-bromo-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 44) and 2-cyanobenzyl zinc bromide. MS (ES+): 372 (base, M+H).

cis-(4a,9b)-5-methyl-8-(3-methyl-butyl)-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole

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The title compound was prepared by following the general coupling procedure as a colorless oil from *cis*-(4a,9b)-8-bromo-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 44) and 3-methylbutyl zinc bromide. MS (ES+): 327 (base, M+H).

EXAMPLE 127

cis-(4a,9b)-4-(5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl)-butyronitrile

The title compound was prepared by following the general coupling procedure as a colorless oil from *cis*-(4a,9b)-8-bromo-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 44) and 3-cyanopropyl zinc bromide. MS (ES+): 324 (base, M+H).

cis-(4a,9b)-8-isobutyl-5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole

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The title compound was prepared by following the general coupling procedure as a colorless oil from *cis*-(4a,9b)-8-bromo-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 44) and isobutyl zinc bromide. MS (ES+): 313 (base, M+H).

EXAMPLE 129

15 (4aS,9bR)-8-isobutyl-5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole

The title compound was prepared by following the general coupling procedure as a colorless oil from (4aS,9bR)-8-bromo-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 45) and *iso*-butyl zinc bromide. MS (ES+): 313 (base, M+H).

cis-(4a,9b)-8-tert-butyl-5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole

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The title compound was prepared by following the general coupling procedure as a colorless oil from *cis*-(4a,9b)-8-bromo-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-10 hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 44) and tert-butyl zinc bromide. MS (ES+): 313 (base, M+H).

EXAMPLE 131

15 cis-(4a,9b)-8-(1-ethyl-propyl)-5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole

The title compound was prepared by following the general coupling procedure as a colorless oil from *cis*-(4a,9b)-8-bromo-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 44) and 1-ethylpropyl zinc bromide. MS (ES+): 327 (base, M+H).

cis-(4a,9b)-5-methyl-8-propyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole

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The title compound was prepared by following the general coupling procedure as a colorless oil from *cis*-(4a,9b)-8-bromo-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-10 hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 44) and propyl zinc bromide. MS (ES+): 299 (base, M+H).

EXAMPLE 133

cis-(4a,9b)-8-butyl-5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole

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The title compound was prepared by following the general coupling procedure as a colorless oil from *cis*-(4a,9b)-8-bromo-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 44) and n-butyl zinc bromide. MS (ES+): 313 (base, M+H).

(4aS,9bR)-8-butyl-5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole

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The title compound was prepared by following the general coupling procedure as a colorless oil from (4aS,9bR)-8-bromo-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-10 hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 45) and n-butyl zinc bromide. MS (ES+): 313 (base, M+H).

EXAMPLE 135

cis-(4a,9b)- 5-methyl-8-pentyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole

The title compound was prepared by following the general coupling procedure as a colorless oil from *cis*-(4a,9b)-8-bromo-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 44) and n-pentyl zinc bromide. MS (ES+): 327 (base, M+H).

cis-(4a,9b)-3-(5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-ylmethyl)-benzonitrile

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The title compound was prepared by following the general coupling procedure as a colorless oil from *cis*-(4a,9b)-8-bromo-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-10 hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 44) and 3-cyanobenzyl zinc bromide. MS (ES+): 372 (base, M+H).

EXAMPLE 137

cis-(4a,9b)-5-methyl-8-phenethyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole

The title compound was prepared by following the general coupling procedure as a colorless oil from *cis*-(4a,9b)-8-bromo-5-methyl-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 44) and phenethyl zinc bromide. MS (ES+): 361 (base, M+H).

cis-(4a,9b)-8-(2-ethyl-butyl)-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole

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The title compound was prepared by following the general coupling procedure as a colorless oil from *cis*-(4a,9b)-8-bromo-6-trifluoromethyl-1,3,4,4a,5,9b-

hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 30) and 2-ethylbutyl zinc bromide. MS (ES+): 327 (base, M+H).

EXAMPLE 139

cis-(4a,9b)-8-(3-methyl-butyl)-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole

The title compound was prepared by following the general coupling procedure as a colorless oil from *cis*-(4a,9b)-8-bromo-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 30) and 3-methylbutyl zinc bromide. MS (ES+): 313 (base, M+H).

cis-(4a,9b)-8-benzyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole

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The title compound was prepared by following the general coupling procedure as a colorless oil from *cis*-(4a,9b)-8-bromo-6-trifluoromethyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 30) and benzyl zinc bromide. MS (ES+): 333 (base, M+H).

EXAMPLE 141

cis-(4a,9b)-8-fluoro-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole

Step A. In a microwave-compatible tube, a solution of (4-fluoro-2-trifluoromethyl-phenyl)-hydrazine monohydrogen chloride (100 mg, 0.43 mmol) and 4-piperidone (67 mg, 0.44 mmol) in 2-PrOH (1.5 mL) was saturated with HCl gas and then sealed. The reaction mixture was irradiated in a microwave at 140 °C for 10 min. The reaction was cooled to 0 °C and filtered. The solid was washed with ether to provide the indole HCl salt (75 mg, 55%) as an off-white solid. A solution of the indole HCl salt (75 mg) in water (10 mL) and CH₂Cl₂ (10 mL) was made basic (pH 10) using K₂CO₃. The basic solution was extracted with CH₂Cl₂ (10 mL) and the

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organic layer was dried over Na₂SO₄, filtered, and evaporated. The residue was triturated with hexanes to provide 8-fluoro-6-trifluoromethyl-2,3,4,5-tetrahydro-1H-pyrido[4,3-b]indole (67 mg, 99%) as a white solid: 1 H NMR (300 MHz, CD₃OD) 8 7.29 (d, J = 10.2 Hz, 1H), 7.11–7.07 (m, 1H), 3.99–3.96 (m, 2H), 3.18–3.14 (m, 2H), 2.87–2.83 (m, 2H); 19 F NMR (282 MHz, CD₃OD) 8 –61.1, –125.4; MS (APCI) 259 (base, M+H)

Step B. Trifluoroacetic acid (5 mL) was added slowly to 8-fluoro-6-trifluoromethyl-2,3,4,5-tetrahydro-1H-pyrido[4,3-b]indole (95 mg, 0.43 mmol) under nitrogen at –10 °C. After 20 min, NaBH₃CN (82 mg, 1.32 mmol) was added over 5 min and the reaction was warmed to room temperature. The reaction was quenched with water (5 mL) and 2 N HCl (5 mL). The reaction mixture was cooled to 0 °C and was made basic (pH 10) using K₂CO₃. The mixture was extracted with CH₂Cl₂ and the combined organic extracts were dried over Na₂SO₄, filtered, and concentrated to provide the crude indoline. The crude indoline was purified by preparative HPLC (Varian Dynamax C18 column, 50–100%, CH₃CN/H₂O with 0.05% TFA) and the residue (34 mg) was dissolved in ether and treated with 1 N HCl to give the title compound (26 mg, 20%) as a white foam: ¹H NMR (300 MHz, CD₃OD) δ 7.30–7.28 (m, 1H), 7.10–7.15 (m, 1H), 4.10–4.05 (m, 1H), 3.53–3.30 (m, 4H), 2.97–2.88 (m, 1H), 2.23–2.18 (m, 2H); ¹⁹F NMR (282 MHz, CD₃OD δ –61.9, –125.2; MS (APCl) 261 (base, M+H).

EXAMPLE 142

cis-(4a,9b)-8-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole

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Step A. A microwave-compatible sealable tube was charged with (4-methyl-2-trifluoromethyl-phenyl)-hydrazine hydrochloride (434 mg, 1.6 mmol), 4-piperidone monohydrate hydrochloride (300 mg, 1.6 mmol), and 2-PrOH (3 mL). The reaction was saturated with HCl gas and the tube was sealed. The reaction mixture was subjected to microwave irradiation at 120 °C for 12 min. The solids were filtered and washed with ether to provide crude 8-methyl-6-trifluoromethyl-2,3,4,5-tetrahydro-1H-pyrido[4,3-b]indole hydrochloride. The resulting crude product was dissolved in THF (12 mL) and H₂O (3 mL), then Na₂CO₃ (184 mg, 1.7 mmol) and Boc₂O (374 mg, 1.7 mmol) were added consecutively. After 2 h, EtOAc (25 mL) and water (10 mL) were added. The aqueous layer was extracted with EtOAc (25 mL) and the combined organic extracts were washed with brine, dried over Na₂SO₄, filtered, and concentrated *in vacuo*. The residue was purified by column chromatography (silica gel, 1–2.5% MeOH/CH₂Cl₂) to provide 8-methyl-6-trifluoromethyl-1,3,4,5-tetrahydro-pyrido[4,3-b]indole-2-carboxylic acid tert-butyl ester (200 mg, 29%).

Step B. 8-Methyl-6-trifluoromethyl-1,3,4,5-tetrahydro-pyrido[4,3-b]indole-2-carboxylic acid tert-butyl ester (110 mg, 0.31 mmol) was dissolved in TFA (5 mL) at 0 °C, then NaBH3CN (49 mg, 0.78 mmol) was added. The reaction mixture was stirred for 1 h at room temperature then quenched with 2 N HCl (2.5 mL). The mixture was made basic (pH = 9) with 6 N NaOH and extracted with EtOAc (2 × 25 mL). The combined organic layers were washed with H_2O (25 mL) and brine (25 mL), dried over Na2SO4, filtered, and concentrated *in vacuo*. Purification of the residue by column chromatography [silica gel, 5–50% (80:18:2 CHCl₃/MeOH/concd NH4OH)/CH₂Cl₂] provided the title compound (44 mg, 37%) as a light yellow oil: 1H NMR (300 MHz, CDCl₃) δ 7.15–7.18 (m, 2H), 4.04–3.92 (m, 1H), 3.80–3.74 (m, 1H), 3.20–3.05 (m, 2H), 3.01–2.75 (m, 3H), 2.17 (s, 3H), 1.98–1.82 (m, 2H), 1.76–1.63 (m, 1H); ESI MS m/z 257 [C₁₃H₁₅F₃N₂ + H]⁺.

cis-(4a,9b)-8-methoxy-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole

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Step A. A microwave-compatible seal tube was charged with (4-methoxy-2-trifluoromethyl-phenyl)-hydrazine hydrochloride (406 mg, 1.7 mmol), 4-piperidone monohydrate hydrochloride (268 mg, 1.7 mmol), and 2-PrOH (4 mL). The reaction mixture was saturated with HCl gas and the tube was sealed. The reaction mixture was subjected to microwave irradiation at 120 °C for 12 min. The solids were filtered, washed with ether and treated with sat. NaHCO₃ (10 mL). The basic solution was extracted with EtOAc (2 × 25 mL) and the combined organic extracts were dried over Na₂SO₄, filtered, and concentrated in vacuo. Purification of the residue by column chromatography [silica gel, 5–75% (80:18:2 CHCl₃/MeOH/concd NH₄OH)/CH₂Cl₂] provided 8-methoxy-6-trifluoromethyl-2,3,4,5-tetrahydro-1H-pyrido[4,3-b]indole (192 mg, in 42%) as an off-white solid: mp 140–144 °C; ¹H NMR (300 MHz, CD₃OD) δ 7.10 (d, J = 2.0 Hz, 1H), 6.93 (d, J = 2.1 Hz, 1H), 3.97 (s, 2H), 3.83 (s, 3H), 3.16 (t, J = 5.8 Hz, 2H), 2.87 (t, J = 5.7 Hz, 2H); ¹⁹F NMR (282 MHz, CD₃OD) δ -61.0; ESI MS 271 [C₁₃H₁₃F₃N₂O + H]⁺.

Step B. To a solution of 8-methoxy-6-trifluoromethyl-2,3,4,5-tetrahydro-1H-pyrido[4,3-b]indole (77 mg, 0.28 mmol) in TFA (4 mL) at 0 °C was added NaBH3CN (43 mg, 0.68 mmol), and the reaction was stirred for 2 h. The reaction was quenched with 2 N HCl (2 mL) at rt then made basic (pH = 9) with 6 N NaOH. The aqueous layer was extracted with EtOAc (2 × 15 mL) and the combined organic layers were washed with brine (15 mL), dried over Na₂SO₄, filtered, and concentrated in vacuo. Purification of the residue by column chromatography [silica gel, 5–50% (80:18:2 CHCl₃/MeOH/concd NH₄OH)/CH₂Cl₂] provided the title compound (49 mg, 63%)

as an off-white semisolid: ¹H NMR (300 MHz, CD₃OD) δ 6.96 (d, J = 2.3 Hz, 1H), 6.73 (d, 1H, J = 2.5 Hz, 1H), 3.94–3.87 (m, 1H), 3.74 (s, 3H), 3.12–2.86 (m, 3H), 2.80–2.61 (m, 2H), 1.93–1.69 (m, 2H); ¹⁹F NMR (282 MHz, CD₃OD) δ –61.3; ESI MS 273 [C₁₃H₁₅F₃N₂O + H]⁺.

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EXAMPLE 144

cis-(4a,9b)-8-bromo-6-cyano-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid tert-butyl ester

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Step A. To a solution of 2-iodophenyl hydrazine hydrochloride (1.0 eq.) and 4-piperidone (1.0 eq.) in 2,2,2-trifluoroethanol was added 12 N HCl (2.0 eq.). The reaction mixture was heated at 60–65 °C for 3h and cooled to rt. then filtered. The residue was purified by column chromatography to provide 6-iodo-2,3,4,5-tetrahydro-1H-pyrido[4,3-b]indole in 69% yield.

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Step B. To a solution of 6-iodo-2,3,4,5-tetrahydro-1H-pyrido[4,3-b]indole in THF and H₂O (3 mL) was added Na₂CO₃ (1.1 eq.) and Boc₂O (1.1 eq.),

consecutively. After 2 h, EtOAc and water were added. The aqueous layer was extracted with EtOAc (25 mL) and the combined organic extracts were washed with brine, dried over Na₂SO₄, filtered, and concentrated *in vacuo*. The residue was purified by column chromatography to give 6-iodo-1,3,4,5-tetrahydro-pyrido[4,3-b]indole-2-carboxylic acid tert-butyl ester in 88% yield.

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Step C. 6-Iodo-2,3,4,5-tetrahydro-1H-pyrido[4,3-b]indole (7.70 g, 25 mmol), tris(dibenzylideneacetone)dipalladium(0) (482 mg, 0.52 mmol), zinc (202 mg, 3.0 mmol), Zn(CN)₂ (1.819 g, 15 mmol), and 1,1'-(diphenylphosphino)ferrocene (572 mg, 1.03 mmol) were dissolved in *N*,*N*-dimethylacetamide (40 mL). The reaction was heated at 120 °C for 1h then cooled to room temperature. The reaction mixture was

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diluted with CH₂Cl₂, filtered through diatomaceous earth, and concentrated. The crude material was dissolved in THF (15 mL) and water (3mL) and treated with K₂CO₃ (3.74 g, 27 mmol) and (Boc)₂O (5.91 g, 27 mmol). After 2 h, the reaction was diluted with water and extracted with CH₂Cl₂ (4 × 20 mL). The combined CH₂Cl₂ layers were dried over Na₂SO₄, filtered, and concentrated. The crude material was purified by flash column chromatography (silica gel, 5–25% EtOAc/hexanes) to afford 6-cyano-1,3,4,5-tetrahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (5.24 g, 91%) as an oil.

Step D. A solution of 6-cyano-1,3,4,5-tetrahydro-pyrido[4,3-b]indole-2-carboxylic acid tert-butyl ester (1.00 g) in CH₂Cl₂ (10 mL) was treated with TFA (2 mL). The reaction mixture was stirred for 7 h then converted to the free base with K₂CO₃ to provide 2,3,4,5-tetrahydro-1H-pyrido[4,3-b]indole-6-carbonitrile (0.66 g, 99%) as a white solid: mp 230–235 °C; 1 H NMR (300 MHz, CD₃OD) δ 7.71–7.69 (m, 1H), 7.50–7.46 (m, 1H), 7.16–7.11 (m, 1H), 4.05–4.04 (m, 2H), 3.26–3.21 (m, 2H), 2.91–2.87 (m, 2H); ESI MS m/z 198 [C₁₂H₁₁N₃ + H]⁺.

Step E. Trifluoroacetic acid (15 mL) was added slowly to 2,3,4,5-tetrahydro-1H-pyrido[4,3-b]indole-6-carbonitrile (900 mg, 4.56 mmol) under nitrogen at -10 °C. After 20 min, sodium cyanoborohydride (860 mg, 13.68 mmol) was added over 20 min and the reaction was stirred for 2 h at -10 °C. The reaction was quenched with water and the solution was made basic (pH 9) using K2CO3. The mixture was extracted with CH2Cl2 (2 x 4 L), dried over Na2SO4, filtered, and concentrated to provide the crude indoline. To a stirred solution of indoline in THF (15 mL) and H₂O (1 mL) was added K2CO3 (662 mg, 4.79 mmol) and Boc2O (1.05 g, 4.79 mmol) at room temperature. After 1 h, the reaction mixture was diluted with EtOAc (1 L) and the layers were separated. The organic phase was dried over Na2SO4, filtered, and evaporated. The residue was purified by flash column chromatography (silica gel, 5-50% EtOAc/hexanes) to afford cis-(4a,9b)-6-cyano-1,3,4,4a,5,9b-hexahydropyrido[4,3-b]indole-2-carboxylic acid tert-butyl ester (250 mg, 20%) as white solid: ¹H NMR (300 MHz, CDCl₃) δ 7.22–7.18 (m, 2H), 6.68 (d, J = 7.8 Hz, 1H), 4.46– 4.44 (m, 1H), 4.19–4.12 (m, 1H), 3.90–3.50 (m, 4H), 2.08–1.96 (m, 1H), 1.86–1.72 (m, 1H), 1.44 (s, 9H); ESI MS m/z 244 $[C_{17}H_{21}N_3O_2 - C_4H_8 + H]^+$.

Step F. To a solution of *cis*-(4a,9b)-6-cyano-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (693 mg, 2.3 mmol) in DMF (20 mL) at 0 °C was added NBS (436 mg, 2.4 mmol). The reaction mixture was stirred at 0 °C for 1 h then quenched by adding crushed ice and then water. The mixture was extracted with Et₂O (2 × 75 mL) and the combined organic extracts were washed with brine (50 mL), dried over Na₂SO₄, filtered, and evaporated *in vacuo*. Purification of the residue by column chromatography (silica gel, 5–30% Et₂O/hexanes) provided the title compound as a yellow solid (782 mg, 90%). MS (ESI): 378 (Base M+H).

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EXAMPLE 145

cis-(4a,9b)-8-bromo-6-cyano-3,4,4a,9b-tetrahydro-1H-pyrido[4,3-b]indole-2,5-dicarboxylic acid di-tert-butyl ester

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A solution of *cis*-(4a,9b)-8-bromo-6-cyano-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (770 mg, 2.0 mmol). It was dissolved in DMF (25 mL), cooled to 0 °C and NaH (254 mg, 6.3 mmol) was carefully added. After 30 min at 0 °C, Boc₂O (1.01 g, 4.6 mmol) was added. The reaction mixture was warmed to room temperature and stirred for 45 min. The reaction mixture was quenched by adding ice and water (20 mL) and extracted with Et₂O (2 × 50 mL). The combined organic layers were washed with water (50 mL) and brine (50 mL), dried over Na₂SO₄, filtered, and concentrated *in vacuo*. Purification of the residue by column chromatography (silica gel, 5–30% Et₂O/hexanes) provided *cis*-(4a,9b)-8-bromo-6-cyano-3,4,4a,9b-tetrahydro-1H-pyrido[4,3-b]indole-2,5-dicarboxylic acid di-*tert*-butyl ester (880 mg, 92%) as a yellow solid: mp 74–78°C; ¹H NMR (300 MHz, CDCl₃) δ 6.58 (br s, 1H), 7.51 (br s,

1H), 4.74–4.60 (m, 1H), 4.38–4.10 (m, 1H), 3.82–3.32 (m, 3H), 3.17–2.78 (m, 1H), 2.16–2.06 (m, 2H), 1.59 (s, 9H), 1.50–1.37 (m, 9H); ESI MS m/z 478 [C₂₂H₂₈BrN₃O₄ + H]⁺.

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EXAMPLE 146

cis-(4a,9b)-8-bromo-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole-6-carbonitrile

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The title compound was prepared from Example150 by following the procedure for Example 72 as a yellow solid. MS (ESI): 278 (Base, M+H).

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EXAMPLE 147

(4aS,9bR)-8-bron20-6-cyano-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester

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(4aS,9bR)-6-cyano-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester were obtained by Prep. HPLC chiral separation (Chiralpak AD column; 80:20 heptane/2-PrOH) from the racemate Example 150 Step E (1st peak eluted) in >99% ee.

The title compound was prepared by following procedures as described in Step F for Example 150. MS (ESI): 378 (Base M+H).

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EXAMPLE 148

(4aS,9bR)-8-bromo-6-cyano-3,4,4a,9b-tetrahydro-1H-pyrido[4,3-b]indole-2,5-dicarboxylic acid di-tert-butyl ester

The title compound was prepared by following procedures as described for Example 151 from (4aS,9bR)-8-bromo-6-cyano-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester. MS (ESI): 478 (Base M+H).

EXAMPLE 149

cis-(4a,9b)-8-bromo-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole-6-carbonitrile

The title compound was prepared from Example 153 by following the procedure for Example 72 as a yellow solid. MS (ESI): 278 (Base, M+H).

cis-(4a,9b)-8-bromo-6-cyano-5-methyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid tert-butyl ester

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Step A. To a solution of *cis*-(4a,9b)-6-cyano-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 150 step E, 900 mg, 3.0 mmol, 1.0 mole equivalent) in DMF (15 mL) was added NaH (0.24 g, 10 mmol, 3.3 mole equivalent). The reaction mixture was stirred at 0 °C for 10 min before MeI (2 mL, 31 mmol, 10 mole equivalent) was added dropwise. The reaction mixture was warmed to rt and stirred for 0.5 h, quenched with water (100 mL) and extracted with EtOAc (3×50 mL). The organic solution was dried (Na₂SO₄), concentrated in vacuo and the residue was chromatographed on a silica gel column by elution with EtOAc/Hexane (gradient) to give *cis*-(4a,9b)-5-methyl-6-cyamo-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester as a white solid (865 mg, 92%).

Step B. To a solution of *cis*-(4a,9b)-5-methyl-6-cyano-1,3,4,4a,5,9b20 hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (722 mg, 2.3 mmol) in DMF (20 mL) at 0 °C was added NBS (436 mg, 2.4 mmol). The reaction mixture was stirred at 0 °C for 1 h then quenched by adding crushed ice and then water. The mixture was extracted with EtOAc (2 × 75 mL) and the combined organic extracts were washed with brine (50 mL), dried over Na₂SO₄, filtered, and evaporated *in*25 *vacuo*. The crude product was chromatographed to provide *cis*-(4a,9b)-8-bromo-5-methyl-6-cyano-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (731 mg, 81%). MS (ESI): 392 (Base, M+H).

cis-(4a,9b)-8-bromo-5-methyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole-6-carbonitrile

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The title compound was prepared from Example 156 by following the procedure for Example 72 as a yellow solid. MS (ESI): 292 (Base, M+H).

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EXAMPLE 152

(4aS,9bR)-8-bromo-6-cyano-5-methyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester

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The title compound was prepared by following procedures as described in Step A B and for Example 156 from (4aS,9bR)-6-cyano-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester. MS (ESI): 392 (Base, M+H).

(4aS,9bR)-8-bromo-5-methyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole-6-carbonitrile

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The title compound was prepared from Example 158 by following the procedure for Example 72 as a yellow solid. MS (ESI): 292 (Base, M+H).

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EXAMPLE 154

cis-(4a,9b)-8-amino-6-cyano-5-methyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid tert-butyl ester

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The title compound was prepared from Example 156 by following the procedure for Example 89 as a yellow solid. MS (ESI): 329 (Base, M+H).

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cis-(4a,9b)-8-amino-5-methyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole-6-carbonitrile

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cis-(4a,9b)-6-Cyano-5-methyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid tert-butyl ester was dissolved in CH2Cl2/TFA (5/1) at rt stirred for 1
h. Upon concentration in vacuo, the residue was partitioned between CH2Cl2/1 M
NaOH. The aqueous phase was extracted with CH2Cl2. The combined organic phases were dried over MgSO4, filtered, and concentrated in vacuo to obtain the title compound MS (ESI): 229 (base, M+H).

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EXAMPLE 156

(4aS,9bR)-8-amino-6-cyano-5-methyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester

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The title compound was prepared from Example 158 by following the procedure for Example 89 as a yellow solid. MS (ESI): 329 (Base, M+H).

(4aS,9bR)-8-amino-5-methyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole-6-carbonitrile

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The title compound was prepared from Example 162 by following the procedure for Example 161 as a yellow solid. MS (ESI): 229 (Base, M+H).

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EXAMPLE 158

cis-(4a,9b)-8-(2-methoxy-pyridin-3-ylamino)-5-methyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole-6-carbonitrile

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The title compound was prepared by following the general method for preparation of (5-Methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl)-pyridin-3-yl-amine (Method A) as an oil (70 mg, 42%) from *cis*-(4a,9b)-8-bromo-6-cyano-5-methyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 156, 196 mg, 0.5 mmol), 2-methoxy-pyridin-3-ylamine(186 mg, 1.5 mmol) and NaO*t*-Bu (144 mg, 1.5 mmol). MS (ESI): 336 (base, M+H).

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(4aS,9bR)-8-(2-methoxy-pyridin-3-ylamino)-5-methyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole-6-carbonitrile

5

The title compound was prepared by following the general method for preparation of (5-Methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl)-pyridin-3-yl-amine (Method A) from (4aS,9bR)-8-bromo-6-cyano-5-methyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 158, 196 mg, 0.5 mmol). MS (ESI): 336 (base, M+H).

EXAMPLE 160

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(4aS,9bR)-5-methyl-8-(pyridin-3-ylamino)-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole-6-carbonitrile

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The title compound was prepared by following the general method for (5-Methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl)-pyridin-3-yl-amine (Method A) as an oil (40 mg, 26%) from (4aS,9bR)-8-bromo-6-cyano-5-methyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 158, 196 mg, 0.5 mmol), 3-amino-pyridine (141 mg, 1.5 mmol) and NaOt-Bu (144 mg, 1.5 mmol). MS (ESI): 306 (base, M+H).

(4aS,9bR)-8-(2-cyano-pyridin-3-ylamino)-5-methyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole-6-carbonitrile

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The title compound was prepared by following the general method for (5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl)-pyridin-3-yl-amine (Method B) as a yellow solid (35 mg, 35%) from (4aS,9bR)-8-amino-6-cyano-3,4,4a,9b-tetrahydro-1H-pyrido[4,3-b]indole-2,5-dicarboxylic acid *tert*-butyl ester (Example 162, 100 mg, 0.31 mmol), 3-bromo-pyridine-2-carbonitrile (56 mg, 0.31 mmol) and Cs₂CO₃ (199 mg, 0.6 mmol). MS (ESI): 331 (base, M+H).

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EXAMPLE 162

(4aS,9bR)-5-methyl-8-(4-trifluoromethyl-pyridin-3-ylamino)-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole-6-carbonitrile

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The title compound was prepared by following the general method for (5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl)-pyridin-3-yl-amine (Method A) as a yellow solid (50 mg, 26%) from (4aS,9bR)-8-bromo-6-cyano-5-methyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 158, 196 mg, 0.5 mmol), 4-trifluoromethyl-pyridin-3-

ylamine (141 mg, 1.5 mmol) and NaOt-Bu (144 mg, 1.5 mmol). MS (ESI): 374 (base, M+H).

EXAMPLE 163

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(4aS,9bR)-5-methyl-8-(6-trifluoromethyl-pyridin-3-ylamino)-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole-6-carbonitrile

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The title compound was prepared by following the general method for (5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl)-pyridin-3-yl-amine (Method A) as an oil (25 mg, 13%) from (4aS,9bR)-8-bromo-6-cyano-5-methyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 158, 196 mg, 0.5 mmol), 6-trifluoromethyl-pyridin-3-ylamine (141 mg, 1.5 mmol) and NaO*t*-Bu (144 mg, 1.5 mmol). MS (ESI): 374 (base, M+H).

EXAMPLE 164

20 (4aS,9bR)-8-(2-ethoxy-pyridin-3-ylamino)-5-methyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole-6-carbonitrile

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The title compound was prepared by following the general method for (5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl)-pyridin-3-yl-amine (Method B) as a yellow solid (30 mg, 28%) from (4aS,9bR)-8-

amino-6-cyano-3,4,4a,9b-tetrahydro-1H-pyrido[4,3-b]indole-2,5-dicarboxylic acid *tert*-butyl ester (Example 162, 100 mg, 0.31mmol), 3-bromo-2-ethoxypyridine (63 mg, 0.31 mmol) and Cs₂CO₃ (199 mg, 0.6 mmol). MS (ESI): 350 (base, M+H).

EXAMPLE 165

(4aS,9bR)-8-(2-isopropoxy-pyridin-3-ylamino)-5-methyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole-6-carbonitrile

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The title compound was prepared by following the general method for (5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl)-pyridin-3-yl-amine (Method B) as a tan solid (32 mg, 21%) from (4aS,9bR)-8-amino-6-cyano-3,4,4a,9b-tetrahydro-1H-pyrido[4,3-b]indole-2,5-dicarboxylic acid *tert*-butyl ester (Example 162, 150 mg, 0.46mmol), 3-bromo-2-*iso*-propoxypyridine (91 mg, 0.42 mmol) and NaOt-Bu (66 mg, 0.69 mmol). ¹H NMR (300 MHz, CDCl₃) δ 7.58 (dd, J = 5.0, 1.4 Hz, 1H), 7.18–6.97 (m, 3H), 6.71 (dd, J = 7.6, 5.0 Hz, 1H), 5.83 (br s, 1H), 5.41 (sept, J = 6.2 Hz, 1H), 3.53–3.48 (m, 1H), 3.33–3.01 (m, 4H), 2.99–2.80 (m, 2H), 2.79–2.61 (m, 1H), 2.59–2.01 (m, 1H), 2.00–1.80 (m, 2H), 1.39 (d, J = 6.2 Hz, 6H); APCI MS m/z 364 [C₂₁H₂₅N₅O + H]⁺.

(4aS,9bR)-8-(6-chloro-pyridin-3-ylamino)-5-methyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole-6-carbonitrile

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The title compound was prepared by following the general method for (5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl)-pyridin-3-yl-amine (Method A) as an oil (20 mg, 12%) from (4aS,9bR)-8-bromo-6-cyano-5-methyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 158, 196 mg, 0.5 mmol), 6-chloro-pyridin-3-ylamine(186 mg, 1.5 mmol) and NaOt-Bu (144 mg, 1.5 mmol). MS (ESI): 340 (base, M+H).

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EXAMPLE 167

(4aS,9bR)-8-(2,5-dichloro-pyridin-3-ylamino)-5-methyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole-6-carbonitrile

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The title compound was prepared by following the general method for (5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl)-pyridin-3-yl-amine (Method A) as a yellow solid (60 mg, 32%) from (4aS,9bR)-8-bromo-6-cyano-5-methyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 158, 196 mg, 0.5 mmol), 2,5-dichloro-pyridin-3-

ylamine(186 mg, 1.5 mmol) and NaOt-Bu (144 mg, 1.5 mmol). MS (ESI): 375 (base, M+H).

EXAMPLE 168

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(4aS,9bR)-8-(6-fluoro-5-methyl-pyridin-3-ylamino)-5-methyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole-6-carbonitrile

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The title compound was prepared by following the general method for (5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl)-pyridin-3-yl-amine (Method A) as an oil (50 mg, 30%) from (4aS,9bR)-8-bromo-6-cyano-5-methyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 158, 196 mg, 0.5 mmol), 6-fluror-5-methyl-pyridin-3-ylamine(186 mg, 1.5 mmol) and NaOt-Bu (144 mg, 1.5 mmol). MS (ESI): 338 (base, M+H).

EXAMPLE 169

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(4aS,9bR)-8-(2-methoxy-5-methyl-pyridin-3-ylamino)-5-methyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole-6-carbonitrile

25

The title compound was prepared by following the general method for (5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl)-

pyridin-3-yl-amine (Method B) as a tan solid (57 mg, 39%) from (4aS,9bR)-8-amino-6-cyano-3,4,4a,9b-tetrahydro-1H-pyrido[4,3-b]indole-2,5-dicarboxylic acid *tert*-butyl ester (Example 162, 150 mg, 0.46mmol), 3-bromo-2-methoxy-5-methylpyridine (84 mg, 0.42 mmol) and NaO*t*-Bu (66 mg, 0.69 mmol): mp 76–80 °C; 1 H NMR (300 MHz, CDCl₃) δ 7.38 (m, 1H), 7.07 (d, J = 2.2 Hz, 1H), 7.00 (d, J = 1.8 Hz, 1H), 6.91 (d, J = 1.8 Hz, 1H), 5.79 (br s, 1H), 3.99 (s, 3H), 3.63–3.51 (m, 1H), 3.23–3.00 (m, 4H), 2.98–2.79 (m, 2H), 2.73–2.26 (m, 2H), 2.18 (s, 3H), 1.99–1.81 (m, 2H); ESI MS m/z 350 [C₂₀H₂₃N₅O + H]⁺.

10 **EXAMPLE 170**

(4aS,9bR)-8-(2-chloro-6-trifluoromethyl-pyridin-3-ylamino)-5-methyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole-6-carbonitrile

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The title compound was prepared by following the general method for (5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl)-pyridin-3-yl-amine (Method A) as an oil (40 mg, 20%) from (4aS,9bR)-8-bromo-6-cyano-5-methyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 158, 196 mg, 0.5 mmol), 2-chloro-6-trifluoromethyl-pyridin-3-ylamine(186 mg, 1.5 mmol) and NaOt-Bu (144 mg, 1.5 mmol). MS (ESI): 408 (base, M+H).

(4aS,9bR)-8-(2,6-dichloro-4-trifluoromethyl-pyridin-3-ylamino)-5-methyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole-6-carbonitrile

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The title compound was prepared by following the general method for (5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl)-pyridin-3-yl-amine (Method A) as an oil (35mg, 16%) from (4aS,9bR)-8-bromo-6-cyano-5-methyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 158, 196 mg, 0.5 mmol), 2,6-dichloro-4-trifluoromethyl-pyridin-3-ylamine(186 mg, 1.5 mmol) and NaO*t*-Bu (144 mg, 1.5 mmol). MS (ESI): 443 (base, M+H).

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General method for preparation of 8-arylamino-5-methyl-6-cyano-2,3,4,4a,5,9bhexahydro-1H-pyrido[4,3-b]indole

8-Bromo-5-methyl-6-cyano-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (1.0 eq.), 2,4-dimethoxyaniline (1.33 eq.), and NaO*t*-Bu (2.4 eq.) were dissolved in anhydrous toluene (0.06 M) while stirring under an argon atmosphere in a sealable test tube. The mixture was degassed with argon for 30 min. Tris(dibenzylideneacetone)dipalladium(0) (0.01 eq.) and 2,2'-bis(diphenylphosphino)-1,1'-binaphthyl (0.03 eq.) were added; the reaction was sealed and heated at 85 °C for 16 h. The reaction was cooled to room temperature, diluted with EtOAc, filtered through a bilayer pad of diatomaceous earth and silica gel, and concentrated *in vacuo*. Purification of the residue by flash column chromatography (silica gel, 10–50% EtOAc/hexanes) provided 6-cyano-8-arylamino-5-methyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid tert-butyl ester. The intermediate was dissolved in CH₂Cl₂ / TFA (4/1) at 0 °C. The reaction mixture was stirred for 3 h then basified with K₂CO₃. The solution was extracted

with CH₂Cl₂ then the organic layer was dried over Na₂SO₄, filtered, and concentrated in vacuo. The crude material was purified by reverse phase prep. HPLC to afford 8-arylamino-5-methyl-6-cyano-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole TFA salt.

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EXAMPLE 172

(4aS,9bR)-8-(2-methoxy-phenylamino)-5-methyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole-6-carbonitrile

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Following the general procedure described above, the title compound was prepared (67 mg, 62%) as a yellow solid using (4aS,9bR)-8-bromo-6-cyano-5-methyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example158, 100 mg, 0.25 mmol), 2-methoxy-aniline (40 mg, 0.33 mmol), and NaO*t*-Bu (56 mg, 0.59 mmol): ¹H NMR (300 MHz, DMSO-*d*₆) δ 8.58 (br s, 1H), 7.30–7.22 (m, 2H), 7.00–6.95 (m, 3H), 6.83–6.80 (m, 2H), 3.80 (s, 3H), 3.70–3.28 (m, 4H), 3.18–3.09 (m, 1H), 3.00 (m, 3H), 2.78–2.69 (m, 1H), 2.10–1.97 (m, 2H); APCI MS *m*/*z* 335 [C₂₀H₂₂N₄O + H]⁺.

EXAMPLE 173

(4aS,9bR)-5-methyl-8-(2-trifluoromethoxy-phenylamino)-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole-6-carbonitrile

Following the general procedure described above, the title compound was prepared (15 mg, 12%) as a tan solid using (4aS,9bR)-8-bromo-6-cyano-5-methyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example158, 100 mg, 0.25 mmol), 2-trifluoromethoxyaniline (58 mg, 0.33 mmol), and NaO*t*-Bu (56 mg, 0.59 mmol): 1 H NMR (300 MHz, DMSO- 2 d6) δ 8.59 (br s, 1H), 7.80 (s, 1H), 7.29–7.26 (m, 2H), 7.18–7.16 (m, 1H), 7.07–7.03 (m, 2H), 6.88–6.85 (m, 1H), 3.91–3.29 (m, 4H), 3.18–2.98 (m, 4H), 2.88–2.72 (m, 1H), 2.10–2.01 (m, 2H); APCI MS 2 d7 (C20H₁₉F₃N₄O + H]⁺.

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EXAMPLE 174

(4aS,9bR)-8-(2-ethoxy-phenylamino)-5-methyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole-6-carbonitrile

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Following the general procedure described above, the title compound was prepared (15 mg, 13%) as a yellow-orange solid using (4aS,9bR)-8-bromo-6-cyano-5-methyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example158, 100 mg, 0.25 mmol), 2-ethoxy-aniline (45 mg, 0.33 mmol), and NaO*t*-Bu (56 mg, 0.59 mmol): 1 H NMR (300 MHz, DMSO- 2 d6) 5 8 8.58 (br s, 1H), 7.26 (s, 1H), 6.98–6.92 (m, 3H), 6.81–6.78 (m, 2H), 4.03 (q, 2 J = 6.8 Hz, 2H), 3.85–3.28 (m, 5H), 3.18–2.98 (m, 4H), 2.80–2.70 (m, 1H), 2.13–2.00 (m, 2H), 1.31 (t, 2 J = 6.9 Hz, 3H); APCI MS 2 M/z 349 [C₂₁H₂₄N₄O + H]⁺.

(4aS,9bR)-8-(2-fluoro-phenylamino)-5-methyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole-6-carbonitrile

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Following the general procedure described above, the title compound was prepared (45 mg, 43%) as a light green solid as a yellow solid using (4aS,9bR)-8-bromo-6-cyano-5-methyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example158, 100 mg, 0.25 mmol), 2-fluoroaniline (36 mg, 0.33 mmol), and NaO*t*-Bu (56 mg, 0.59 mmol): 1 H NMR (300 MHz, DMSO- 2 d6) δ 8.60 (br s, 1H), 7.80 (br s, 1H), 7.24–7.23 (m, 1H), 7.20–6.98 (m, 3H), 6.91–6.90 (m 1H), 6.88–6.80 (m, 1H), 3.61–3.28 (m, 4H), 3.14–2.97 (m, 4H), 2.80–2.70 (m, 1H), 2.10–1.98 (m, 2H); ESI MS 2 323 [C₁₉H₁₉FN₄ + H]⁺.

EXAMPLE 176

(4aS,9bR)-5-methyl-8-o-tolylamino-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole-6-carbonitrile

Following the general procedure described above, the title compound was prepared (67 mg, 62%) as a yellow solid using (4aS,9bR)-8-bromo-6-cyano-5-methyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example158, 100 mg, 0.25 mmol), 2-methylaniline (35 mg, 0.33 mmol), and NaOt-

Bu (56 mg, 0.59 mmol): 1 H NMR (300 MHz, DMSO- d_{6}) δ 9.60 (br s, 1H), 7.23–7.12 (m, 3H), 7.10–7.03 (m, 1H), 6.98–6.93 (m, 1H), 6.84–6.80 (m, 2H), 3.61–3.33 (m, 4H), 3.18–3.09 (m, 1H), 3.01 (s, 3H), 2.78–2.70 (m, 1H), 2.20–1.93 (m, 5H); APCI MS m/z 319 $[C_{20}H_{22}N_{4} + H]^{+}$.

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EXAMPLE 177

(4aS,9bR)-8-(2-ethyl-phenylamino)-5-methyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole-6-carbonitrile

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Following the general procedure described above, the title compound was prepared (70 mg, 63%) as a light green solid using (4aS,9bR)-8-bromo-6-cyano-5-methyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example158, 100 mg, 0.25 mmol), 2-ethylaniline (40 mg, 0.33 mmol), and NaOt-Bu (56 mg, 0.59 mmol): 1 H NMR (300 MHz, DMSO- d_{6}) δ 8.63 (br s, 1H), 7.34–7.17 (m, 3H), 7.11–7.05 (m, 1H), 6.99–6.96 (m, 1H), 6.92–6.87 (m, 1H). 6.81–6.80 (m, 1H), 4.20–3.91 (m, 1H), 3.55–3.49 (m, 1H), 3.39–3.28 (m, 2H), 3.13–3.09 (m, 1H), 3.02–3.00 (m, 3H), 2.75–2.49 (m, 3H), 2.14–1.93 (m, 2H), 1.14 (t, J = 7.5 Hz, 3H); APCI MS m/z 333 [C_{21} H $_{24}$ N $_{4}$ + H] $_{-}^{+}$.

(4aS,9bR)-8-(4-fluoro-phenylamino)-5-methyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole-6-carbonitrile

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Following the general procedure described above, the title compound was prepared (49 mg, 47%) as a light green solid using (4aS,9bR)-8-bromo-6-cyano-5-methyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example158, 100 mg, 0.25 mmol), 4-fluoroaniline (36 mg, 0.33 mmol), and NaO*t*-Bu (56 mg, 0.59 mmol): ¹H NMR (300 MHz, DMSO-*d*₆) δ 8.59 (br s, 1H), 7.90 (br s, 1H), 7.25–7.24 (m, 1H), 7.08–7.01 (m, 2H), 6.95–6.91 (m, 3H), 3.68–3.30 (m, 4H), 3.18–2.99 (m, 4H), 2.80–2.72 (m, 1H), 2.11–1.99 (m, 2H); APCI MS *m/z* 323 [C₁₉H₁₉FN₄ + H]⁺.

EXAMPLE 179

(4aS,9bR)-5-methyl-8-m-tolylamino-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole-6-carbonitrile

Following the general procedure described above, the title compound was prepared (61 mg, 58%) as a yellow solid using (4aS,9bR)-8-bromo-6-cyano-5-methyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example158, 100 mg, 0.25 mmol), 3-methylaniline (35 mg, 0.33 mmol), and NaOt-

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Bu (56 mg, 0.59 mmol): ¹H NMR (300 MHz, DMSO- d_6) δ 8.55 (br s, 1H), 7.90 (br s, 1H), 7.30–7.29 (m, 1H), 7.07 (t, J = 7.5 Hz, 1H), 6.97–6.96 (m, 1H), 6.72–6.70 (m, 2H), 6.60 (d, J = 7.5 Hz, 1H), 3.60–3.29 (m, 4H), 3.20–3.11 (m, 1H), 3.04 (s, 3H), 2.81–2.73 (m, 1H), 2.21 (s, 3H), 2.12–1.97 (m, 2H); APCI MS m/z 319 [C₂₀H₂₂N₄ + H]⁺.

EXAMPLE 180

(4aS,9bR)-8-(2-cyano-phenylamino)-5-methyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole-6-carbonitrile

Following the general procedure described above, the title compound was prepared (7 mg, 10%) as a yellow solid using (4aS,9bR)-8-bromo-6-cyano-5-methyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example158, 62 mg, 0.116 mmol), 2-cyanoaniline (24 mg, 0.21 mmol), and Cs₂CO₃ (118 mg, 0.36 mmol): ¹H NMR (300 MHz, DMSO-*d*₆) δ 8.59 (br s, 1H), 8.31 (s, 1H), 7.60–7.58 (m, 1H), 7.48–7.40 (m, 1H), 7.33–7.32 (m, 1H), 7.10–7.09 (m, 1H), 7.00–6.98 (m, 1H), 6.90–6.84 (m, 1H), 3.72–3.29 (m, 4H), 3.12–3.00 (m, 4H), 2.88–2.80 (m, 1H), 2.04–2.00 (m, 2H); ESI MS *m/z* 330 [C₂₀H₁₉N₅ + H]⁺.

(4aS,9bR)-8-(2,4-dimethoxy-phenylamino)-5-methyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole-6-carbonitrile

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Following the general procedure described above, the title compound was prepared (25 mg, 21%) as a yellow solid using (4aS,9bR)-8-bromo-6-cyano-5-methyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example158, 100 mg, 0.25 mmol), 2,4-dimethoxyaniline (51 mg, 0.33 mmol), and NaO*t*-Bu (56 mg, 0.59 mmol): ¹H NMR (300 MHz, DMSO-*d*₆) δ 6.96–6.90 (m, 3H), 6.62–6.51 (m, 1H), 6.43–6.39 (m, 1H), 5.51 (br s, 1H), 3.84 (s, 3H), 3.79 (s, 3H), 3.48–3.45 (m, 1H), 3.13–3.02 (m, 5H), 2.89–2.86 (m, 2H), 2.67–2.61 (m, 1H), 1.90–1.87 (m, 2H); APCI MS *m/z* 365 [C₂₁H₂₄N₄O₂ + H]⁺.

EXAMPLE 182

(4aS,9bR)-8-(2,3-difluoro-phenylamino)-5-methyl-2,3,4,4a,5,9b-hexahydro-1 H-pyrido [4,3-b] indole-6-carbonitrile

Following the general procedure described above, the title compound was prepared (12 mg, 11%) as a yellow solid using (4aS,9bR)-8-bromo-6-cyano-5-methyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example158, 100 mg, 0.25 mmol), 2,3-difluoro-aniline (42 mg, 0.33 mmol), and

NaO*t*-Bu (56 mg, 0.59 mmol): ¹H NMR (300 MHz, DMSO-*d*₆) δ 7.09–7.08 (m, 1H), 7.04–7.03 (m 1H), 6.92–6.89 (m, 1H), 6.70–6.59 (m, 2H), 5.62 (br, 1H), 3.61–3.59 (m, 1H), 3.34–3.29 (m, 1H), 3.18–3.11 (m, 4H), 3.03–2.90 (m, 2H), 2.71–2.63 (m, 1H), 2.11–1.82 (m, 2H); APCI MS m/z 341 [C₁₉H₁₈F₂N₄ + H]⁺.

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EXAMPLE 183

(4aS,9bR)-8-(5-fluoro-2-methyl-phenylamino)-5-methyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole-6-carbonitrile

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Following the general procedure described above, the title compound was prepared (28 mg, 25%) as a yellow solid using (4aS,9bR)-8-bromo-6-cyano-5-methyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example158, 100 mg, 0.25 mmol), 5-fluoro-2-methylaniline (41 mg, 0.33 mmol), and NaO*t*-Bu (56 mg, 0.59 mmol): ¹H NMR (300 MHz, CDCl₃) δ 7.07–6.98 (m, 3H), 6.55–6.45 (m, 2H), 5.20 (br s, 1H), 3.57–3.55 (m, 1H), 3.18–3.08 (m, 5H), 2.92–2.88 (m, 2H), 2.72–2.68 (m, 1H), 2.18–1.92 (m, 5H); APCI MS *m/z* 337 [C₂₀H₂₁FN₄ + 40 H]⁺.

EXAMPLE 184

(4aS,9bR)-8-(4-fluoro-3-methyl-phenylamino)-5-methyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole-6-carbonitrile

Following the general procedure described above, the title compound was prepared (25 mg, 23%) as a yellow solid using (4aS,9bR)-8-bromo-6-cyano-5-methyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example158, 100 mg, 0.25 mmol), 4-fluoro-3-methylaniline (41 mg, 0.33 mmol), and NaO*t*-Bu (56 mg, 0.59 mmol): 1 H NMR (300 MHz, CDCl3, DMSO- 2 d6) 5 9.46 (br s, 1H), 7.09–7.08 (m, 1H), 7.05–7.04 (m, 1H), 6.97–6.73 (m, 3H), 3.61–3.29 (m, 4H), 3.18–2.91 (m, 3H), 2.69–2.58 (m, 2H), 2.37–2.16 (m, 5H); APCI MS 2 d7 (C₂₀H₂₁FN₄ + H]⁺.

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EXAMPLE 185

(4aS,9bR)-8-(4-chloro-3-methyl-phenylamino)-5-methyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole-6-carbonitrile

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(4aS,9bR)-8-(4-fluoro-2-methyl-phenylamino)-5-methyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole-6-carbonitrile

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Following the general procedure described above, the title compound was prepared (8 mg, 7%) as a tan solid using (4aS,9bR)-8-bromo-6-cyano-5-methyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example158, 100 mg, 0.25 mmol), 4-fluoro-2-methylaniline (41 mg, 0.33 mmol), and NaO*t*-Bu (56 mg, 0.59 mmol):): 1 H NMR (300 MHz, DMSO- 2 d6) δ 8.65 (br s, 1H), 7.24 (s, 1H), 7.10–6.91 (m, 3H), 6.71–6.70 (m, 1H), 6.59 (br s, 1H), 3.55–3.30 (m, 4H), 3.20–2.98 (m, 4H), 2.77–2.68 (m, 1H), 2.21–1.92 (m, 5H); APCI MS 2 d7 (2 20H21FN4 + H] $^{+}$.

EXAMPLE 187

(4aS,9bR)-8-(4-chloro-2-methyl-phenylamino)-5-methyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole-6-carbonitrile

Following the general procedure described above, the title compound was prepared (75 mg, 65%) as a light green solid using (4aS,9bR)-8-bromo-6-cyano-5-methyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example 158, 100 mg, 0.25 mmol), 4-chloro-2-methylaniline (46 mg, 0.33 mmol),

and NaOt-Bu (56 mg, 0.59 mmol):): ¹H NMR (300 MHz, DMSO-t6) δ 8.58 (br s, 1H), 7.28 (s, 1H), 7.19–7.18 (m, 2H), 7.10–7.07 (m, 1H), 6.92–6.90 (m, 2H), 3.80–3.52 (m, 2H), 3.42–3.31 (m, 2H), 3.20–3.09 (m, 1H), 3.02 (s, 3H), 2.79–2.70 (m, 1H), 2.20–1.97 (m, 5H); APCI MS t7/z 353 [C₂₀H₂₁ClN₄ + H]⁺.

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EXAMPLE 188

(4aS,9bR)-8-(2,4-difluoro-phenylamino)-5-methyl-2,3,4,4a,5,9b-hexahydro-1 H-pyrido [4,3-b] indole-6-carbonitrile

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Following the general procedure described above, the title compound was prepared (52 mg, 47%) as a light green solid using (4aS,9bR)-8-bromo-6-cyano-5-methyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example158, 100 mg, 0.25 mmol), 2,4-difluoro-aniline (42 mg, 0.33 mmol), and NaO*t*-Bu (56 mg, 0.59 mmol):): 1 H NMR (300 MHz, DMSO- 2 d $_{0}$) δ 8.57 (br s, 1H), 7.78 (s, 1H), 7.29–7.06 (m, 3H), 6.92–6.89 (m, 1H), 6.84 (s, 1H), 3.89–3.48 (m, 2H), 3.38–3.29 (m, 2H), 3.19–3.07 (m, 1H), 3.01 (s, 3H), 2.73–2.69 (m, 1H), 2.10–1.98 (m, 2H); APCI MS 2 M/z 341 [2 C₁₉H₁₈F₂N₄ + H]⁺.

EXAMPLE 189

(4aS,9bR)-8-(3,4-difluoro-phenylamino)-5-methyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole-6-carbonitrile

Following the general procedure described above, the title compound was prepared (52 mg, 47%) as a light green solid using (4aS,9bR)-8-bromo-6-cyano-5-methyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example158, 100 mg, 0.25 mmol), 3,4-difluoro-aniline (42 mg, 0.33 mmol), and NaO*t*-Bu (56 mg, 0.59 mmol):): 1 H NMR (300 MHz, DMSO- 2 d₆) δ 8.60 (br s, 1H), 8.10 (s, 1H), 7.31–7.18 (m, 2H), 7.00–6.98 (m, 1H), 6.89–6.80 (m, 1H), 6.70–6.63 (m, 1H), 4.03–3.58 (m, 2H), 3.49–3.32 (m, 2H), 3.20–3.00 (m, 4H), 2.89–2.78 (m, 1H), 2.10–2.01 (m, 2H); APCI MS 2 d₇ 341 [C₁₉H₁₈F₂N₄ + H]⁺.

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EXAMPLE 190

(4aS,9bR)-8-(3-chloro-4-fluoro-phenylamino)-5-methyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole-6-carbonitrile

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Following the general procedure described above, the title compound was prepared (53 mg, 44%) as a light green solid using (4aS,9bR)-8-bromo-6-cyano-5-methyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example158, 100 mg, 0.25 mmol), 3-chloro-4-fluoroaniline (48 mg, 0.33 mmol), and NaO*t*-Bu (56 mg, 0.59 mmol):): 1 H NMR (300 MHz, DMSO- 2 d6) δ 8.59 (br s, 1H), 8.10–8.06 (m, 1H), 7.33–7.19 (m, 2H), 7.03–6.94 (m, 2H), 6.87–6.84 (m, 1H), 3.78–3.30 (m, 4H), 3.19–3.00 (m, 4H), 2.85–2.77 (m, 1H), 2.09–1.99 (m, 2H); APCI MS 2 m/z 357 [C₁₉H₁₈ClFN₄ + H]⁺.

(4aS,9bR)-8-(2,4-dimethyl-phenylamino)-5-methyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole-6-carbonitrile

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Following the general procedure described above, the title compound was prepared (60 mg, 54%) as a yellow solid using (4aS,9bR)-8-bromo-6-cyano-5-methyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example158, 100 mg, 0.25 mmol), 2,4-dimethyl-aniline (44 mg, 0.33 mmol), and NaO*t*-Bu (56 mg, 0.59 mmol):): ¹H NMR (300 MHz, DMSO-*d*₆) δ 8.64 (br s, 1H), 7.18 (br s, 1H), 7.10–7.09 (m, 1H), 7.01–7.00 (m, 1H), 6.91–6.90 (m, 2H), 6.60–6.59 (m, 1H), 4.02–3.82 (m, 1H), 3.53–3.48 (m, 1H), 2.39–3.31 (m, 2H), 3.19–3.11 (m, 1H), 3.00 (s, 3H), 2.74–2.62 (m, 1H), 2.23 (s, 3H), 2.13–1.95 (m, 5H); APCI MS *m/z* 333 [C₂₁H₂₄N₄ + H]⁺.

EXAMPLE 192

20 (4aS,9bR)-8-(4-methoxy-2-methyl-phenylamino)-5-methyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole-6-carbonitrile

Following the general procedure described above, the title compound was prepared (82 mg, 72%) as a light green solid using (4aS,9bR)-8-bromo-6-cyano-5-methyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester

(Example158, 100 mg, 0.25 mmol), 4-methoxy-2-methylaniline (45 mg, 0.33 mmol), and NaOt-Bu (56 mg, 0.59 mmol):): 1 H NMR (300 MHz, DMSO- 2 d6) δ 8.59 (br s, 1H), 7.11 (br s, 1H), 6.97–6.90 (m, 2H), 6.80–6.79 (m, 1H), 6.71–6.70 (m, 1H), 6.51–6.50 (m, 1H), 3.79–3.41 (m, 5H), 3.34–3.25 (m, 2H), 3.19–3.08 (m, 1H), 3.02–2.97 (m, 3H), 2.70–2.62 (m, 1H), 2.12–1.96 (m, 5H); APCI MS m/z 349 [C₂₁H₂₄N₄O + H]⁺.

EXAMPLE 193

10 (4aS,9bR)-8-(5-chloro-2-methoxy-phenylamino)-5-methyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole-6-carbonitrile

Following the general procedure described above, the title compound was prepared (74 mg, 60%) as a light green solid using (4aS,9bR)-8-bromo-6-cyano-5-methyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example158, 100 mg, 0.25 mmol), 5-chloro-2-methoxyaniline (52 mg, 0.33 mmol), and NaO*t*-Bu (56 mg, 0.59 mmol):): ¹H NMR (300 MHz, DMSO-*d*₆) δ 8.59 (br s, 1H), 7.43 (s, 1H), 7.35–7.34 (s, 1H), 7.08–7.07 (s, 1H), 6.96–6.93 (m, 1H), 6.80–6.74 (m, 2H), 3.83–3.52 (m, 5H), 3.48–3.28 (m, 2H), 3.19–2.98 (m, 4H), 2.82–2.71 (m, 1H), 2.18–1.98 (m, 2H); APCI MS *m*/z 369 [C₂₀H₂₁ClN₄O + H]⁺.

(4aS,9bR)-8-(2,4-dichloro-phenylamino)-5-methyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole-6-carbonitrile

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Following the general procedure described above, the title compound was prepared (45 mg, 37%) as a yellow solid using (4aS,9bR)-8-bromo-6-cyano-5-methyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example158, 100 mg, 0.25 mmol), 2,4-dichloro-aniline (53 mg, 0.33 mmol), and NaO*t*-Bu (56 mg, 0.59 mmol):): ¹H NMR (300 MHz, DMSO-*d*₆) δ 8.58 (br s, 1H), 7.63–7.62 (m, 1H), 7.51–7.52 (m, 1H), 7.31–7.30 (m, 1H), 7.20–7.18 (m, 1H), 7.10–7.08 (m, 1H), 6.98–6.97 (m, 1H), 3.80–3.62 (m, 1H), 3.49–3.30 (m, 3H), 3.18–3.01 (m, 4H), 2.88–2.79 (m, 1H), 2.07–1.99 (m, 2H); ESI MS *m/z* 373 [C₁₉H₁₈Cl₂N₄ + H]⁺.

EXAMPLE 195

(4aS,9bR)-8-(2,5-dimethyl-phenylamino)-5-methyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole-6-carbonitrile

Following the general procedure described above, the title compound was prepared (70 mg, 64%) as a yellow solid using (4aS,9bR)-8-bromo-6-cyano-5-methyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example158, 100 mg, 0.25 mmol), 2,5-dimethyl-aniline (53 mg, 0.33 mmol), and

NaOt-Bu (56 mg, 0.59 mmol):): ¹H NMR (300 MHz, DMSO- d_6) δ 8.54 (br s, 1H), 7.15–7.13 (m, 2H), 7.02–7.00 (m, 1H), 6.80–6.75 (m, 2H), 6.66–6.63 (m, 1H), 3.54–3.52 (m, 1H), 3.40–3.30 (m, 3H), 3.19–3.13 (m, 1H), 3.01 (s, 3H), 2.79 (m, 1H), 2.18 (s, 3H), 2.11 (s, 3H), 2.08–1.93 (m, 2H); ESI MS m/z 333 [C₂₁H₂₄N₄ + H]⁺.

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EXAMPLE 196

(4aS,9bR)-8-(2,5-difluoro-phenylamino)-5-methyl-2,3,4,4a,5,9b-hexahydro-1 H-pyrido [4,3-b] indole-6-carbonitrile

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Following the general procedure described above, the title compound was prepared (27 mg, 24%) as a light green solid using (4aS,9bR)-8-bromo-6-cyano-5-methyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example158, 100 mg, 0.25 mmol), 2,5-difluoro-aniline (53 mg, 0.33 mmol), and NaO*t*-Bu (56 mg, 0.59 mmol):): 1 H NMR (300 MHz, DMSO- 2 d6) δ 8.59 (br s, 1H), 8.01 (br s, 1H), 7.38–7.37 (m, 1H), 7.21–7.12 (m, 1H), 7.05–7.04 (m, 1H), 6.69–6.62 (m, 1H), 6.60–6.54 (m, 1H), 3.69–3.31 (m, 4H), 3.19–3.00 (m, 4H), 2.89–2.81 (m, 1H), 2.11–2.05 (m, 2H); ESI MS 2 M 2 341 [C₁₉H₁₈F₂N₄ + H] $^{+}$.

EXAMPLE 197

(4aS,9bR)-8-(2,6-difluoro-phenylamino)-5-methyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole-6-carbonitrile

Following the general procedure described above, the title compound was prepared (10 mg, 9%) as a yellow solid using (4aS,9bR)-8-bromo-6-cyano-5-methyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example158, 100 mg, 0.25 mmol), 2,6-difluoro-aniline (53 mg, 0.33 mmol), and NaO*t*-Bu (56 mg, 0.59 mmol):): 1 H NMR (300 MHz, DMSO- 2 d6) δ .8.59 (br s, 1H), 7.81 (s, 1H), 7.12–7.08 (m, 3H), 6.92–6.90 (m, 1H), 6.58–6.57 (m, 1H), 3.50–3.29 (m, 4H), 3.20–3.11 (m, 1H), 2.98 (s, 3H), 2.70–2.62 (m, 1H), 2.18–1.92 (m, 2H); ESI MS m Z 341 [2 C₁₉H₁₈F₂N₄ + H]⁺.

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EXAMPLE 198

(4aS,9bR)-8-iso-butyl-5-methyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido [4,3-b] indole-6-carbonitrile

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Following the general procedure for Example 127-146, the title compound was prepared (13 mg, 48%) as a light yellow oil using (4aS,9bR)-8-bromo-6-cyano-5-methyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example158, 40 mg, 0.10 mmol), *iso*-butylzinc bromide (0.5 M in THF, 2.5 mole equivalent) and Pd(PPh3)4 (0.06 mole equivalent).: MS (ESI): 270 (base, M+H).

(4aS,9bR)-5-methyl-8-(3-methyl-butyl)-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole-6-carbonitrile

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Following the general procedure for Example 127-146, the title compound was prepared (11 mg, 39%) as a light yellow oil using (4aS,9bR)-8-bromo-6-cyano-5-methyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example158, 40 mg, 0.10 mmol), 3-methyl-butylzinc bromide (0.5 M in THF, 2.5 mole equivalent) and Pd(PPh3)4 (0.06 mole equivalent).: MS (ESI): 284 (base, M+H).

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EXAMPLE 200

(4aS,9bR)-8-butyl-5-methyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b] indole-6-carbonitrile

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Following the general procedure for Example 127-146, the title compound was prepared (8 mg, 30%) as a light yellow oil using (4aS,9bR)-8-bromo-6-cyano-5-methyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example158, 40 mg, 0.10 mmol), n-butylzinc bromide (0.5 M in THF, 2.5 mole equivalent) and Pd(PPh3)4 (0.06 mole equivalent).: MS (ESI): 270 (base, M+H).

(4aS,9bR)-8-benzyl-5-methyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole-6-carbonitrile

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Following the general procedure for Example 127-146, the title compound was prepared (25 mg, 83%) as a light yellow oil using (4aS,9bR)-8-bromo-6-cyano-5-methyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example158, 40 mg, 0.10 mmol), benzylzinc bromide (0.5 M in THF, 2.5 mole equivalent) and Pd(PPh3)4 (0.06 mole equivalent).: MS (ESI): 304 (base, M+H).

EXAMPLE 202

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(4aS,9bR)-8-(2,4-difluoro-benzyl)-5-methyl-2,3,4,4a,5,9b-hexahydro-1 H-pyrido [4,3-b] indole-6-carbonitrile

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Following the general procedure for Example 127-146, the title compound was prepared (20 mg, 59%) as a light yellow oil using (4aS,9bR)-8-bromo-6-cyano-5-methyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example158, 40 mg, 0.10 mmol), 2,4-difluoro-benzylzinc bromide (0.5 M in THF, 2.5 mole equivalent) and Pd(PPh3)4 (0.06 mole equivalent).: MS (ESI): 340 (base, M+H).

(4aS,9bR)-8-(2,5-difluoro-benzyl)-5-methyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole-6-carbonitrile

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Following the general procedure for Examples 127-146, the title compound was prepared (18 mg, 53%) as a light yellow oil using (4aS,9bR)-8-bromo-6-cyano-5-methyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example158, 40 mg, 0.10 mmol), 2,5-difluoro-benzylzinc bromide (0.5 M in THF, 2.5 mole equivalent) and Pd(PPh₃)₄ (0.06 mole equivalent).: MS (ESI): 340 (base, M+H).

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EXAMPLE 204

(4aS,9bR)-5-methyl-8-phenethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole-6-carbonitrile

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Following the general procedure for Examples 127-146, the title compound was prepared (8 mg, 25%) as a light yellow oil using (4aS,9bR)-8-bromo-6-cyano-5-methyl-1,3,4,4a,5,9b-hexahydro-pyrido[4,3-b]indole-2-carboxylic acid *tert*-butyl ester (Example158, 40 mg, 0.10 mmol), phenethylzinc bromide (0.5 M in THF, 2.5 mole equivalent) and Pd(PPh3)4 (0.06 mole equivalent).: MS (ESI): 318 (base, M+H).

cis-(4a,9b)-8-(2-methoxy-phenylamino)-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole-6-carbonitrile

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The title compound was prepared by following the general method for *cis*-(4a,9b)- (5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl)-pyridin-3-yl-amine (Method B) as a yellow solid (35 mg, 24%) from cis-(4a,9b)-8-amino-6-cyano-3,4,4a,9b-tetrahydro-1H-pyrido[4,3-b]indole-2,5-dicarboxylic acid *tert*-butyl ester (Example 151, 187 mg, 0.45mmol), 1-bromo-2-methoxy-benzene (84 mg, 0.45 mmol) and NaOtBu (87 mg, 0.9 mmol). MS (ESI): 321 (base, M+H).

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EXAMPLE 206

(4aS,9bR)-8-(2,6-dimethyl-pyridin-3-ylamino)-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indole-6-carbonitrile

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The title compound was prepared by following the general method for cis-(4a,9b)- (5-methyl-6-trifluoromethyl-2,3,4,4a,5,9b-hexahydro-1H-pyrido[4,3-b]indol-8-yl)-pyridin-3-yl-amine (Method B) as a yellow solid (35 mg, 24%) from cis-(4a,9b)-8-amino-6-cyano-3,4,4a,9b-tetrahydro-1H-pyrido[4,3-b]indole-2,5dicarboxylic acid tert-butyl ester (Example 151, 187 mg, 0.45mmol), 3-bromo-2,6-

PH7483 NP

dimethyl-pyridine (84 mg, 0.45 mmol) and Cs2CO3 (293 mg, 0.9 mmol). MS (ESI): 320 (base, M+H).

TABLE 1

$$R^8$$
 H
 R^6
 R^5

Ex#	R ⁵	R ⁶	R ⁸	b	R1
1	H	MeS-	Br	cis	Н
2	Me	MeS-	Br	cis	Н
7	Me	4-Me-Ph-S-	Br	trans	Me
8	Me	4-Me-Ph-S-	Br	Cis	Me
9	Me	4-Me-Ph-S-	Ph-S-	trans	Me
10	Boc	Me-	Br	Cis	Boc
11	Boc	Me-	Br	4aS,9bR	Boc
12	Boc	Me-	H ₂ N-	4aS,9bR	Boc
24	Н	F ₃ C-	Br	Cis	Boc
25	H	F ₃ C-	Br	4aS,9bR	Boc
38	Me	F ₃ C-	Br	Cis	Boc
39	Me	F ₃ C-	Br	4aS,9bR	Boc
64	Me	F ₃ C-	OH	Cis	Boc
65	Me	F ₃ C-	OH	4aS,9bR	Boc
66	Me	F ₃ C-	OH	Cis	H
67	Me	F ₃ C-	OH	4aS,9bR	Н
83	Me	F ₃ C-	H2N	Cis	Boc
84	Me	F ₃ C-	H2N	4aS,9bR	Boc
85	Me	F ₃ C-	H2N	Cis	H
86	Me	F ₃ C-	H2N	4aS,9bR	Н
141	Н	F ₃ C-	F-	Cis	Н
142	Н	F ₃ C-	Me-	Cis	Н
143	Н	F ₃ C-	OMe-	Cis	Н
144	Н	NC-	Br	Cis	Boc
145	Boc	NC-	Br	Cis	Boc
146	Н	NC-	Br	Cis	Н
147	Н	NC-	Br	4aS,9bR	Boc
148	Boc	NC-	Br	4aS,9bR	Boc
149	Н	NC-	Br	4aS,9bR	H
150	Me	NC-	Br	Cis	Boc
151	Me	NC-	Br	Cis	H

Ex#	R ⁵	R ⁶	R ⁸	b	R1
152	Me	NC-	Br	4aS,9bR	Boc
153	Me	NC-	Br	4aS,9bR	H
154	Me	NC-	H2N-	Cis	Boc
155	Me	NC-	H2N-	Cis	H
156	Me	NC-	H2N-	4aS,9bR	Boc
157	Me	NC-	H2N-	4aS,9bR	H

TABLE 2

Ex#	R ⁵	R ⁶	b	R ⁸
3	Н	MeS-	cis	-NH ₂
4	Н	MeS-	cis	Ph-NH-
5	Н	MeS-	cis	4-F-Ph-NH-
6	Н	MeS-	cis	2-Me-4-MeO-Ph-NH-
13	H	Me-	4aS,9bR	2,3-diCl-Ph-NH-
14	Н	Me-	4aS,9bR	3,4-diCl-Ph-NH-
15	Н	Me-	4aS,9bR	3-Cl-4-Me-Ph-NH-
16	H	Me-	4aS,9bR	2,4-diCl-Ph-NH-
_17	H	Me-	4aS,9bR	2,6-diCl-Ph-NH-
18	Н	Me-	4aS,9bR	2,4-diF-Ph-NH-
19	H	Me-	4aS,9bR	2-MeO-5-Me-Ph-NH-
20	H	Me-	4aS,9bR	3-Cl-4-CN-Ph-NH-
21	H	Me-	4aS,9bR	2-F-4-Cl-Ph-NH-
22	H	Me-	4aS,9bR	2-F-5-CF ₃ -Ph-NH-
23	H	Me-	4aS,9bR	2-Cl-5-CF ₃ -Ph-NH-
26	Н	F ₃ C-	Cis	Cyclohexyl-NH-
27	Н	F ₃ C-	Cis	Me ₂ CH ₂ CH(CH ₃)-NH-
28	H	F ₃ C-	Cis	Benzyl-NH-
29	Н	F ₃ C-	Cis	1-Phenyl-ethyl-NH-
30	Н	F ₃ C-	Cis	2-Me-Benzyl-NH-
31	Н	F ₃ C-	Cis	2-OMe-Benzyl-NH-
32	Н	F ₃ C-	Cis	2-Cl-6-F-Benzyl-NH-
33	Н	F ₃ C-	Cis	4-t-Bu-Benzyl-NH-
34	Н	F ₃ C-	Cis	3-Me-Benzyl-NH-

Ex#	R ³	R ⁶	b	R ⁸
35	Н	F ₃ C-	Cis	4-Me-Benzyl-NH-
36	Н	F ₃ C-	Cis	2,5-diMe-Benzyl-NH-
37	Н	F ₃ C-	Cis	2-F-4-CF ₃ -Benzyl-NH-
40	Me	F ₃ C-	Cis	Cyclohexyl-NH-
41	Me	F ₃ C-	Cis	Benzyl-NH-
42	Me	F ₃ C-	Cis	2-CF ₃ -Benzyl-NH-
43	Me	F ₃ C-	Cis	1-Phenyl-ethyl-NH-
44	Me	F ₃ C-	Cis	(s)1-cyclohexyl-ethyl-NH-
45	Me	F ₃ C-	Cis	Exo-bicyclo[2,2,1] hept-2-yl-NH-
46	Me	F ₃ C-	Cis	(S)-2-Phenyl-propyl-NH-
47	Me	F ₃ C-	4aS,9bR	2-SMe-Ph-NH-
48	Me	F ₃ C-	4aS,9bR	2-Et-Ph-NH-
49	Me	F ₃ C-	4aS,9bR	2-OMe-5-Me-Ph-NH-
50	Me	F ₃ C-	4aS,9bR	Ph-NH-
51	Me	F ₃ C-	4aS,9bR	2-F-Ph-NH-
52	Me	F ₃ C-	4aS,9bR	3-F-Ph-NH-
53	Me	F ₃ C-	4aS,9bR	4-F-Ph-NH-
54	Me	F ₃ C-	4aS,9bR	2-OEt-Ph-NH-
55	Me	F ₃ C-	4aS,9bR	2-Me-Ph-NH-
56	Me	F ₃ C-	4aS,9bR	3-F-4-Me-Ph-NH-
57	Me	F ₃ C-	4aS,9bR	3-Cl-4-F-Ph-NH-
58	Me	F ₃ C-	4aS,9bR	4-F-3-Me-Ph-NH-
59	Me	F ₃ C-	4aS,9bR	4-Cl-3-Me-Ph-NH-
60	Me	F ₃ C-	4aS,9bR	2-OMe-Ph-NH-
61	Me	F ₃ C-	4aS,9bR	2,6-diMe-Ph-NH-
62	Me	F ₃ C-	4aS,9bR	2,4-diF-Ph-NH-
63	Me	F ₃ C-	4aS,9bR	2,6-diF-Ph-NH-
68	Me	F ₃ C-	Cis	Cyclopropylmethoxy-
69	Me	F ₃ C-	Cis	Cyclopentylmethoxy-
70	Me	F ₃ C-	Cis	3-me-butoxy-
71	Me	F ₃ C-	Cis	n-propoxy-
72	Me	F ₃ C-	Cis	n-butoxy-
73	Me	F ₃ C-	Cis	3,3-diMe-butoxy-
74	Me	F ₃ C-	Cis	Cyclobutylmethoxy-
75	Me	F ₃ C-	4aS,9bR	2-Py-methoxy-
76	Me	F ₃ C-	4aS,9bR	3-Py-methoxy-
77	Me	F ₃ C-	4aS,9bR	4-Py-methoxy-
78	Me	F ₃ C-	Cis	2-Me-Ph-O-

Ex#	R^5	R ⁶	b	R ⁸
79	Me	F ₃ C-	Cis	2,5-diMe-Ph-O-
80	Me	F ₃ C-	Cis	2-CN-Ph-O-
81	Me	F ₃ C-	Cis	4-CN-Ph-O-
82	Me	F ₃ C-	Cis	2-OMe-Ph-O-
87	Me	F ₃ C-	Cis	3-Py-NH-
88	Me	F ₃ C-	Cis	3-(2-Cl-Py)- NH-
89	Me	F ₃ C-	Cis	3-(2-CN-Py)- NH-
90	Me	F ₃ C-	4aS,9bR	3-(2-CN-Py)- NH-
91	Me	F ₃ C-	Cis	3-(4-OMe-Py)- NH-
92	Me	F ₃ C-	4aS,9bR	3-(4-OMe-Py)- NH-
93	Me	F ₃ C-	Cis	3-(4-F-Py)- NH-
94	Me	F ₃ C-	4aS,9bR	3-(4-OMe-Py)- NH-
95	Me	F ₃ C-	Cis	3-(4-CN-Py)- NH-
96	Me	F ₃ C-	Cis	3-(5-CN-Py)- NH-
97	Me	F ₃ C-	Cis	3-(5-COOMe-Py)- NH-
98	Me	F ₃ C-	Cis	3-(5-COOEt-Py)- NH-
99	Me	F ₃ C-	4aS,9bR	3-(2-OMe-Py)- NH-
100	Me	F ₃ C-	Cis	3-(4-Cl-Py)- NH-
101	Me	F ₃ C-	4aS,9bR	3-(4-Cl-Py)- NH-
102	Me	F ₃ C-	Cis	3-(2-OEt-Py)- NH-
103	Me	F ₃ C-	Cis	3-(2,4-diOMe-Py)- NH-
104	Me	F ₃ C-	4aS,9bR	3-(2,4-diOMe-Py)- NH-
105	Me	F ₃ C-	Cis	3-(2,4-diCl-Py)- NH-
106	Me	F ₃ C-	Cis	3-(2-CN-6-Me-Py)- NH-
107	Me	F ₃ C-	Cis	3-(4-F-5-Me-Py)- NH-
108	Me	F ₃ C-	4aS,9bR	3-(4-F-5-Me-Py)- NH-
109	Me	F ₃ C-	Cis	3-(2,5-diCl-Py)- NH-
110	Me	F ₃ C-	Cis	3-(2-CN-5-OMe-Py)- NH-
111	Me	F ₃ C-	Cis	3-(2-OMe-4-Me-Py)- NH-
112	Me	F ₃ C-	Cis	3-(2-Me-4-Cl-Py)- NH-
113	Me	F ₃ C-	Cis	3-(2-CN-4-Me-Py)- NH-
114	Me	F ₃ C-	Cis	3-(2,4-diMe-Py)- NH-
115	Me	F ₃ C-	Cis	3-(2-I-PrO-4-Me-Py)- NH-
116	Me	F ₃ C-	Cis	3-(2-EtO-4-Me-Py)- NH-
117	Me	F ₃ C-	Cis	3-(2-EtO-6-Me-Py)- NH-
118	Me	F ₃ C-	Cis	3-(2-EtO-4-Me-Py)- NH-
119	Me	F ₃ C-	Cis	2-(4-Me-Py)- NH-
120	Me	F ₃ C-	Cis	2-(3-Me-4-Br-Py)- NH-

Ex#	R ⁵	R ⁶	b	R ⁸
121	Me	F ₃ C-	Cis	2-Et-butyl-
122	Me	F ₃ C-	Cis	benzyl-
123	Me	F ₃ C-	4aS,9bR	benzyl-
124	Me	F ₃ C-	Cis	Cyclohex-
125	Me	F ₃ C-	Cis	2-CN-benzyl-
126	Me	F ₃ C-	Cis	2-Et-butyl-
127	Me	F ₃ C-	Cis	3-CN-propyl-
128	Me	F ₃ C-	Cis	isobutyl-
129	Me	F ₃ C-	4aS,9bR	isobutyl-
130	Me	F ₃ C-	Cis	tert-butyl-
131	Me	F ₃ C-	Cis	1-Et-propyl-
132	Me	F ₃ C-	Cis	n-propyl-
133	Me	F ₃ C-	Cis	n-butyl-
134	Me	F ₃ C-	4aS,9bR	n-butyl-
135	Me	F ₃ C-	Cis	n-pentyl-
136	Me	F ₃ C-	Cis	3-CN-benzyl-
137	Me	F ₃ C-	Cis	phenethyl-
138	Н	F ₃ C-	Cis	2-Et-butyl-
139	Н	F ₃ C-	Cis	3-Me-butyl-
140	Н	F ₃ C-	Cis	benzyl-
158	Me	NC-	Cis	3-(2-OMe-Py)-NH-
159	Me	NC-	4aS,9bR	3-(2-OMe-Py)-NH-
160	Me	NC-	4aS,9bR	3-Py-NH-
161	Me	NC-	4aS,9bR	3-(2-CN-Py)-NH-
162	Me	NC-	4aS,9bR	3-(6-CF3-Py)-NH-
163	Me	NC-	4aS,9bR	3-(4-CF3-Py)-NH-
164	Me	NC-	4aS,9bR	3-(2-OEt-Py)-NH-
165	Me	NC-	4aS,9bR	3-(2-iOPr-Py)-NH-
166	Me	NC-	4aS,9bR	3-(4-Cl-Py)-NH-
167	Me	NC-	4aS,9bR	3-(2,5-diCl-Py)-NH-
168	Me	NC-	4aS,9bR	3-(4-F-5-Me-Py)-NH-
169	Me	NC-	4aS,9bR	3-(2-OMe-5-Me-Py)-NH-
170	Me	NC-	4aS,9bR	3-(2-Cl-4-CF3-Py)-NH-
171	Me	NC-	4aS,9bR	3-(2,4-diCl-6-CF3-Py)-NH-
172	Me	NC-	4aS,9bR	2-OMe-Ph-NH-
173	Me	NC-	4aS,9bR	2-OCF3-Ph-NH-
174	Me	NC-	4aS,9bR	2-OEt-Ph-NH-
175	Me	NC-	4aS,9bR	2-F-Ph-NH-
176	Me	NC-	4aS,9bR	2-Me-Ph-NH-
177	Me	NC-	4aS,9bR	2-Et-Ph-NH-
178	Me	NC-	4aS,9bR	4-F-Ph-NH-

Ex#	R ⁵	R ⁶	b	R ⁸
179	Me	NC-	4aS,9bR	3-Me-Ph-NH-
180	Me	NC-	4aS,9bR	2-CN-Ph-NH-
181	Me	NC-	4aS,9bR	2,4-diOMe-Ph-NH-
182	Me	NC-	4aS,9bR	2,3-diF-Ph-NH-
183	Me	NC-	4aS,9bR	2-Me-5-F-Ph-NH-
184	Me	NC-	4aS,9bR	3-Me-4-F-Ph-NH-
185	Me	NC-	4aS,9bR	3-Me-4-Cl-Ph-NH-
186	Me	NC-	4aS,9bR	2-Me-4-F-Ph-NH-
187	Me	NC-	4aS,9bR	2-Me-4-Cl-Ph-NH-
188	Me	NC-	4aS,9bR	2,4-diF-Ph-NH-
189	Me	NC-	4aS,9bR	2,4-diF-Ph-NH-
190	Me	NC-	4aS,9bR	3-Cl-4-F-Ph-NH-
191	Me	NC-	4aS,9bR	2,4-diMe-Ph-NH-
192	Me	NC-	4aS,9bR	2-Me-4-OMe-Ph-NH-
193	Me	NC-	4aS,9bR	2-OMe-5-Cl-Ph-NH-
194	Me	NC-	4aS,9bR	2,4-diCl-Ph-NH-
195	Me	NC-	4aS,9bR	2,5-diMe-Ph-NH-
196	Me	NC-	4aS,9bR	2,5-diF-Ph-NH-
197	Me	NC-	4aS,9bR	2,6-diF-Ph-NH-
198	Me	NC-	4aS,9bR	Isobutyl-
199	Me	NC-	4aS,9bR	3-Me-butyl-
200	Me	NC-	4aS,9bR	n-butyl-
201	Me	NC-	4aS,9bR	benzyl-
202	Me	NC-	4aS,9bR	2,4-diF-benzyl-
203	Me	NC-	4aS,9bR	2,5-diF-benzyl-
204	Me	NC-	4aS,9bR	Phenethyl-
205	Н	NC-	Cis	2-OMe-Ph-NH-
206	Н	NC-	Cis	3-(2,4-diMe-Py)-NH-

UTILITIES AND COMBINATIONS

Utilities

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The compounds of the present invention are 5HT_{2C} modulators, and include compounds which are, for example, selective agonists, partial agonists, antagonists or partial antagonists of the 5HT_{2C} receptor. Accordingly, the compounds of the present invention may be useful for the treatment or prevention of diseases and disorders associated with 5HT_{2C} receptor activity. Preferably, compounds of the present invention possess activity as agonists of the 5HT_{2C} receptor, and may be used in the treatment of diseases or disorders associated with the activity of the 5HT_{2C} receptor.

Accordingly, the compounds of the present invention can be administered to mammals, preferably humans, for the treatment of a variety of conditions and

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disorders, including, but not limited to metabolic and eating disorders as well as conditions associated with metabolic disorders, (e.g., obesity, diabetes, arteriosclerosis, hypertension, polycystic ovary disease, cardiovascular disease, osteoarthritis, dermatological disorders, impaired glucose hemostatsis, insulin resistance, hypercholesterolemia, hypertriglyceridemia, cholelithiasis and sleep disorders, dislipidemic conditions, bulimia nervosa and compulsive eating disorders); pain; sleep disorders and psychiatric disorders, such as substance abuse, depression, anxiety, psychosis, mania and schizophrenia.

These compounds could also be used for the improvement of cognitive function (e.g., the treatment of dementia, including Alzheimer's disease, short term memory loss and attention deficit disorders); neurodegenerative disorders (e.g., Parkinson's Disease, cerebral apoplexy and craniocerebral trauma) and hypotension (e.g., hemorrhagic and endotoxin-inducd hypotension). These compounds could also be used for treatment of cardiac dysfunction (e.g., associated with valvular disease, myocardial infarction, cardiac hypertrophy or congestive heart failure); and improvement of the overall pulmonary function; transplant rejection; rheumatoid arthritis; osteoarthritis; fibromyalgia; multiple sclerosis; inflammatory bowel disease; lupus; graft vs. host disease; T-cell mediated hypersensitivity disease; psoriasis; asthma; Hashimoto's thyroiditis; Guillain-Barre syndrome; cancer; contact dermatitis; allergic rhinitis; and ischemic or reperfusion injury. These compounds could also be used for treatment of sexual dysfunction and erectogenesis.

Compounds useful in the treatment of appetite or motivational disorders regulate desires to consume sugars, carbohydrates, alcohol or drugs and more generally to regulate the consumption of ingredients with hedonic value. In the present description and in the claims, appetite disorders are understood as meaning: disorders associated with a substance and especially abuse of a substance and/or dependency on a substance, disorders of eating behaviors, especially those liable to cause excess weight, irrespective of its origin, for example: bulimia nervosa, craving for sugars. The present invention therefore further relates to the use of a 5HT_{2C} receptor agonist for the treatment of bulimia and obesity, including obesity associated with type II diabetes (non-insulin-dependent diabetes), or more generally any disease resulting in the patient becoming overweight. Overweight and obesity, as described

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herein, is defined by a body mass index (kg/m²) for example, at least 26. It may be due to any cause, whether genetic or environmental, including overeating and bulemia, polycycstic ovary disease, craniopharyngeoma, Prader-Willi Syndrome, Frohlich's Syndrome, Type II diabetes, growth hormone deficiency, Turner's

5 Syndrome and other pathological states characterized by reduced metabolic activity or reduced energy expenditure. As used with reference to the utilities described herein, the term "treating" or "treatment" encompasses prevention, partial alleviation, or cure of the disease or disorder. Further, treatment of obesity is expected to prevent progression of medical covariants of obesity, such as arteriosclerosis, Type II diabetes, polycystic ovary disease, cardiovascular disease, osteoarthritis, dermatological disorders, hypertension, insulin resistance, hypercholesterolemia, hypertriglyceridemia, cholelithiasis and sleep disorders.

Compounds in the present invention may also be useful in treating substance abuse disorders, including substance dependence or abuse without physiological dependence. Substances of abuse include alcohol, amphetamines (or amphetamine-like substances), caffeine, cannabis, cocaine, hallucinogens, inhalents, nicotine, opioids, phencyclidine (or phencyclidine-like compounds), sedative-hypnotics or benzodiazepines, and other (or unknown) substances and combinations of the above. The terms "substance abuse disorders" also includes drug, nicotine or alcohol withdrawal syndromes and substance-induced anxiety or mood disorder with onset during withdrawal.

Compounds in the present invention may be useful in treating memory impairment and cognitive disorders. The condition of memory impairment is manifested by impairment of the ability to learn new information and/or the inability to recall previously learned information. Memory impairment is a primary symptom of dementia and can also be a symptom associated with such diseases as Alzheimer's disease, schizophrenia, Parkinson's disease, Huntington's disease, Pick's disease, Creutzfeld-Jakob disease, attention deficit-hyperactivity disorder, HIV, cardiovascular disease such as ischemia or stroke, and head trauma as well as age-related cognitive decline. Dementias are diseases that include memory loss and additional intellectual impairment separate from memory. 5HT_{2C} modulators may also be useful in treating

cognitive impairments related to attentional deficits, such as attention deficithyperactivity disorders.

Compounds in the present invention may also be useful in treating diseases associated with dysfunction of brain dopaminergic systems, such as Parkinson's Disease and substance abuse disorders. Parkinsons's Disease is a neurodenerative movement disorder characterized by bradykinesia and tremor.

Combinations

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The present invention includes within its scope pharmaceutical compositions comprising, as an active ingredient, a therapeutically effective amount of at least one of the compounds of formula I, alone or in combination with a pharmaceutical carrier or diluent. Optionally, compounds of the present invention can be used alone, in combination with other suitable therapeutic agents useful in the treatment of the aforementioned disorders including: anti-obesity agents; anti-diabetic agents, appetite suppressants; cholesterol/lipid-lowering agents, cognition enhancing agents, agents used to treat neurodegeneration, agents used to treat respiratory conditions, agents used to treat bowel disorders, anti-inflammatory agents; anti-anxiety agents; anti-depressants; anti-psychotic agents; sedatives; hypnotics; anti-hypertensive agents; anti-tumor agents and analgesics.

Such other therapeutic agent(s) may be administered prior to, simultaneously with, or following the administration of the $5HT_{2C}$ modulators in accordance with the invention.

Examples of suitable anti-obesity agents for use in combination with the compounds of the present invention include leptin and leptin-sensitizing agents, melanocortin receptor (MC4R) agonists, agouti-related peptide (AGRP) antagonists, melanin-concentrating hormone receptor (MCHR) antagonists, growth hormone secretagogue receptor (GHSR) antagonists, orexin antagonists, CCK agonists, GLP-1 agonists, NPY1 or NPY5 antagonsits, NPY2 modulators, corticotropin releasing factor agonists, histamine receptor-3 (H3) modulators, aP2 inhibitors, PPAR gamma modulators, PPAR delta modulators, beta 3 adrenergic agonists, such as AJ9677 (Takeda/Dainippon), L750355 (Merck), or CP331648 (Pfizer) or other known beta 3 agonists as disclosed in U.S. Patent Nos. 5,541,204, 5,770,615, 5,491,134, 5,776,983

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and 5,488,064, a thyroid receptor beta modulator, such as a thyroid receptor ligand as disclosed in WO 97/21993 (U. Cal SF), WO 99/00353 (KaroBio) and GB98/284425 (KaroBio), a lipase inhibitor, such as orlistat or ATL-962 (Alizyme), leptinergics, adiponectin modulating agents, cannabinoid-1 receptor antagonists, such as SR-141716 (Sanofi) or SLV-319 (Solvay) and monoamine reuptake inhibitors or releasing agents, such as fenfluramine, dexfenfluramine, fluvoxamine, fluoxetine, paroxetine, sertraline, chlorphentermine, cloforex, clortermine, picilorex, sibutramine, dexamphetamine, phentermine, phenylpropanolamine or mazindol, anorectic agents such as topiramate (Johnson & Johnson), axokine (Regeneron).

Examples of suitable anti-diabetic agents for use in combination with the compounds of the present invention include: insulin, which may include short- and long-lasting forms as well as oral and inhaled forms, insulin secretagogues or insulin sensitizers, which may include biguanides, sulfonyl ureas, glucosidase inhibitors, aldose reductase inhibitors, PPAR γ agonists such as thiazolidinediones, PPAR α agonists (such as fibric acid derivatives), PPAR δ antagonists or agonists, PPAR α/γ dual agonists such as the compounds described in Bristol-Myers Squibb U.S. patent 6,414,002, dipeptidyl peptidase IV (DP4) inhibitors such as the compounds described in Bristol-Myers Squibb U.S. patents 6,395,767 and 6,573,287, SGLT2 inhibitors such as the compounds described in Bristol-Myers Squibb U.S. patents 6,414,126 and 6,515,117, glycogen phosphorylase inhibitors, and/or meglitinides, as well as insulin, and/or glucagon-like peptide-1 (GLP-1), and/or a PTP-1B inhibitor (protein tyrosine phosphatase-1B inhibitor).

The antidiabetic agent may be glucokinase inhibitors, 11 β HSD inhibitors or oral antihyperglycemic agents, which is preferably a biguanide such as metformin or phenformin or salts thereof, preferably metformin HCl. Where the antidiabetic agent is a biguanide, the compounds of the present invention will be employed in a weight ratio to biguanide within the range from about 0.001:1 to about 10:1, preferably from about 0.01:1 to about 5:1.

The antidiabetic agent may also preferably be a sulfonyl urea such as glyburide (also known as glibenclamide), glimepiride (disclosed in U.S. Patent No. 4,379,785), glipizide, gliclazide or chlorpropamide, other known sulfonylureas or other antihyperglycemic agents which act on the ATP-dependent channel of the beta-

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cells, with glyburide and glipizide being preferred, which may be administered in the same or in separate oral dosage forms. The oral antidiabetic agent may also be a glucosidase inhibitor such as acarbose (disclosed in U.S. Patent No. 4,904,769) or miglitol (disclosed in U.S. Patent No. 4,639,436), which may be administered in the same or in a separate oral dosage forms.

The compounds of the present invention may be employed in combination with a PPAR γ agonist such as a thiazolidinedione oral anti-diabetic agent or other insulin sensitizers (which has an insulin sensitivity effect in NIDDM patients) such as troglitazone (Warner-Lambert's REZULIN, disclosed in U.S. Patent No. 4,572,912), rosiglitazone (SKB), pioglitazone (Takeda), Mitsubishi's MCC-555 (disclosed in U.S. Patent No. 5,594,016), Glaxo-Welcome's GL-262570, englitazone (CP-68722, Pfizer) or darglitazone (CP-86325, Pfizer, isaglitazone (MIT/J&J), JTT-501 (JPNT/P&U), L-895645 (Merck), R-119702 (Sankyo/WL), NN-2344 (Dr. Reddy/NN), or YM-440 (Yamanouchi), preferably rosiglitazone and pioglitazone.

15 The compounds of the present invention may be employed in combination with anti-hyperlipidemia agents, or agents used to treat arteriosclerosis. An example of an hypolipidemic agent would be an HMG CoA reductase inhibitor which includes, but is not limited to, mevastatin and related compounds as disclosed in U.S. Patent No. 3,983,140, lovastatin (mevinolin) and related compounds as disclosed in U.S. 20 Patent No. 4,231,938, pravastatin and related compounds such as disclosed in U.S. Patent No. 4,346,227, simvastatin and related compounds as disclosed in U.S. Patent Nos. 4,448,784 and 4,450,171. Other HMG CoA reductase inhibitors which may be employed herein include, but are not limited to, fluvastatin, disclosed in U.S. Patent No. 5,354,772, cerivastatin disclosed in U.S. Patent Nos. 5,006,530 and 5,177,080, 25 atorvastatin disclosed in U.S. Patent Nos. 4,681,893, 5,273,995, 5,385,929 and 5,686,104, pitavastatin (Nissan/Sankyo's nisvastatin (NK-104) or itavastatin), disclosed in U.S. Patent No. 5,011,930, Shionogi-Astra/Zeneca rosuvastatin (visastatin (ZD-4522)) disclosed in U.S. Patent No. 5,260,440, and related statin compounds disclosed in U.S. Patent No. 5,753,675, pyrazole analogs of 30 mevalonolactone derivatives as disclosed in U.S. Patent No. 4,613,610, indene analogs of mevalonolactone derivatives as disclosed in PCT application WO 86/03488, 6-[2-(substituted-pyrrol-1-yl)-alkyl)pyran-2-ones and derivatives thereof as

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disclosed in U.S. Patent No. 4,647,576, Searle's SC-45355 (a 3-substituted pentanedioic acid derivative) dichloroacetate, imidazole analogs of mevalonolactone as disclosed in PCT application WO 86/07054, 3-carboxy-2-hydroxy-propane-phosphonic acid derivatives as disclosed in French Patent No. 2,596,393, 2,3-disubstituted pyrrole, furan and thiophene derivatives as disclosed in European Patent Application No. 0221025, naphthyl analogs of mevalonolactone as disclosed in U.S. Patent No. 4,686,237, octahydronaphthalenes such as disclosed in U.S. Patent No. 4,499,289, keto analogs of mevinolin (lovastatin) as disclosed in European Patent Application No.0,142,146 A2, and quinoline and pyridine derivatives disclosed in U.S. Patent Nos. 5,506,219 and 5,691,322. In addition, phosphinic acid compounds useful in inhibiting HMG CoA reductase suitable for use herein are disclosed in GB 2205837.

The squalene synthetase inhibitors suitable for use herein include, but are not limited to, α-phosphono-sulfonates disclosed in U.S. Patent No. 5,712,396, those disclosed by Biller et al, J. Med. Chem., 1988, Vol. 31, No. 10, pp 1869-1871, including isoprenoid (phosphinyl-methyl)phosphonates as well as other known squalene synthetase inhibitors, for example, as disclosed in U.S. Patent No. 4,871,721 and 4,924,024 and in Biller, S.A., Neuenschwander, K., Ponpipom, M.M., and Poulter, C.D., Current Pharmaceutical Design, 2, 1-40 (1996).

In addition, other squalene synthetase inhibitors suitable for use herein include the terpenoid pyrophosphates disclosed by P. Ortiz de Montellano et al, J. Med. Chem., 1977, 20, 243-249, the farnesyl diphosphate analog A and presqualene pyrophosphate (PSQ-PP) analogs as disclosed by Corey and Volante, J. Am. Chem. Soc., 1976, 98, 1291-1293, phosphinylphosphonates reported by McClard, R.W. et al, J.A.C.S., 1987, 109, 5544, cyclopropanes reported by Capson, T.L., PhD dissertation, June, 1987, Dept. Med. Chem. U of Utah, Abstract, Table of Contents, pp 16, 17, 40-43, 48-51, Summary, pyrrolidine derivatives as disclosed by Sasyou, et al, WO 02/083636 and N-aryl-substituted cyclic amine derivatives disclosed by Okada et al, WO 02/076973.

Other hypolipidemic agents suitable for use herein include, but are not limited to, fibric acid derivatives, α PPAR agonists, such as fenofibrate, gemfibrozil, clofibrate, bezafibrate, ciprofibrate, clinofibrate and the like, probucol, and related

compounds as disclosed in U.S. Patent No. 3,674,836, probucol, phenylfibrate and gemfibrozil being preferred, bile acid sequestrants such as cholestyramine, colestipol and DEAE-Sephadex (SECHOLEX, POLICEXIDE) and cholestagel (Sankyo/Geltex), as well as lipostabil (Rhone-Poulenc), Eisai E-5050 (an N-500 substituted ethanolamine derivative), imanixil (HOE-402), tetrahydrolipstatin (THL), istigmastanylphos-phorylcholine (SPC, Roche), aminocyclodextrin (Tanabe Seiyoku), Ajinomoto AJ-814 (azulene derivative), melinamide (Sumitomo), Sandoz 58-035, American Cyanamid CL-277,082 and CL-283,546 (disubstituted urea derivatives), nicotinic acid (niacin), acipimox, acifran, neomycin, p-aminosalicylic acid, aspirin, poly(diallylmethylamine) derivatives such as disclosed in U.S. Patent No. 4,759,923, quaternary amine poly(diallyldimethylammonium chloride) and ionenes such as disclosed in U.S. Patent No. 4,027,009, and other known serum cholesterol lowering agents.

The other hypolipidemic agent may be an ACAT inhibitor (which also has 15 anti-atherosclerosis activity) such as disclosed in, Drugs of the Future 24, 9-15 (1999), (Avasimibe); "The ACAT inhibitor, Cl-1011 is effective in the prevention and regression of aortic fatty streak area in hamsters", Nicolosi et al, Atherosclerosis (Shannon, Irel). (1998), 137(1), 77-85; "The pharmacological profile of FCE 27677: a novel ACAT inhibitor with potent hypolipidemic activity mediated by selective 20 suppression of the hepatic secretion of ApoB100-containing lipoprotein", Ghiselli, Giancarlo, Cardiovasc. Drug Rev. (1998), 16(1), 16-30; "RP 73163: a bioavailable alkylsulfinyl-diphenylimidazole ACAT inhibitor", Smith, C., et al, Bioorg. Med. Chem. Lett. (1996), 6(1), 47-50; "ACAT inhibitors: physiologic mechanisms for hypolipidemic and anti-atherosclerotic activities in experimental animals", Krause et 25 al, Editor(s): Ruffolo, Robert R., Jr.; Hollinger, Mannfred A., Inflammation: Mediators Pathways (1995), 173-98, Publisher: CRC, Boca Raton, Fla.; "ACAT inhibitors: potential anti-atherosclerotic agents", Sliskovic et al, Curr. Med. Chem. (1994), 1(3), 204-25; "Inhibitors of acyl-CoA: cholesterol O-acyl transferase (ACAT) as hypocholesterolemic agents. 6. The first water-soluble ACAT inhibitor with lipidregulating activity. Inhibitors of acyl-CoA:cholesterol acyltransferase (ACAT). 7. 30 Development of a series of substituted N-phenyl-N'-[(1phenylcyclopentyl)methyl]ureas with enhanced hypocholesterolemic activity", Stout

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et al, Chemtracts: Org. Chem. (1995), 8(6), 359-62, or TS-962 (Taisho Pharmaceutical Co. Ltd), as well as F-1394, CS-505, F-12511, HL-004, K-10085 and YIC-C8-434.

The hypolipidemic agent may be an upregulator of LDL receptor activity such as MD-700 (Taisho Pharmaceutical Co. Ltd) and LY295427 (Eli Lilly). The hypolipidemic agent may be a cholesterol absorption inhibitor preferably Schering-Plough's SCH48461 (ezetimibe) as well as those disclosed in Atherosclerosis 115, 45-63 (1995) and J. Med. Chem. 41, 973 (1998).

The other lipid agent or lipid-modulating agent may be a cholesteryl transfer protein inhibitor (CETP) such as Pfizer's Torcetrapib® as well as those disclosed in WO/0038722 and in EP 818448 (Bayer) and EP 992496, and Pharmacia's SC-744 and SC-795, as well as CETi-1 and JTT-705.

The hypolipidemic agent may be an ileal Na⁺/bile acid cotransporter inhibitor such as disclosed in Drugs of the Future, 24, 425-430 (1999). The ATP citrate lyase inhibitor which may be employed in the combination of the invention may include, for example, those disclosed in U.S. Patent No. 5,447,954.

The other lipid agent also includes a phytoestrogen compound such as disclosed in WO 00/30665 including isolated soy bean protein, soy protein concentrate or soy flour as well as an isoflavone such as genistein, daidzein, glycitein or equal, or phytosterols, phytostanol or tocotrienol as disclosed in WO 2000/015201; a beta-lactam cholesterol absorption inhibitor such as disclosed in EP 675714; an HDL upregulator such as an LXR agonist, a PPAR α-agonist and/or an FXR agonist; an LDL catabolism promoter such as disclosed in EP 1022272; a sodium-proton exchange inhibitor such as disclosed in DE 19622222; an LDL-receptor inducer or a steroidal glycoside such as disclosed in U.S. Patent No. 5,698,527 and GB 2304106; an anti-oxidant such as beta-carotene, ascorbic acid, α-tocopherol or retinol as disclosed in WO 94/15592 as well as Vitamin C and an antihomocysteine agent such as folic acid, a folate, Vitamin B6, Vitamin B12 and Vitamin E; isoniazid as disclosed in WO 97/35576; a cholesterol absorption inhibitor, an HMG-CoA synthase inhibitor, or a lanosterol demethylase inhibitor as disclosed in WO 97/48701; a PPAR δ agonist for treating dyslipidemia; or a sterol regulating element binding protein-I (SREBP-1) as disclosed in WO 2000/050574, for example, a sphingolipid, such as ceramide, or

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neutral sphingomyelenase (N-SMase) or fragment thereof, and inhibitors or lipid synthesis enzymes such as, for example, ACC, FAS, DGAT, MGAT, GPAT, AMP kinase, CPT1 and SCD1. Preferred dislipidemic agents are pravastatin, lovastatin, simvastatin, atorvastatin, fluvastatin, pitavastatin, rosuvastatin, phenylfibrate and Pfizer's Torcetrapib® as well as niacin and/or cholestagel.

The compounds of the present invention may be employed in combination with anti-hypertensive agents. Examples of suitable anti-hypertensive agents for use in combination with the compounds of the present invention include beta adrenergic blockers, calcium channel blockers (L-type and T-type; e.g. diltiazem, verapamil, nifedipine, amlodipine and mybefradil), diuretics (e.g., chlorothiazide, hydrochlorothiazide, flumethiazide, hydroflumethiazide, bendroflumethiazide, methylchlorothiazide, trichloromethiazide, polythiazide, benzthiazide, ethacrynic acid tricrynafen, chlorthalidone, furosemide, musolimine, bumetanide, triamtrenene, amiloride, spironolactone), renin inhibitors, ACE inhibitors (e.g., captopril, zofenopril, fosinopril, enalapril, ceranopril, cilazopril, delapril, pentopril, quinapril, ramipril, lisinopril), AT-1 receptor antagonists (e.g., losartan, irbesartan, valsartan, candasartan and talmisartan), ET receptor antagonists (e.g., sitaxsentan, atrsentan and compounds disclosed in U.S. Patent Nos. 5,612,359 and 6,043,265), Dual ET/AII antagonist (e.g., compounds disclosed in WO 00/01389), neutral endopeptidase (NEP) inhibitors, vasopepsidase inhibitors (dual NEP-ACE inhibitors) (e.g., omapatrilat and gemopatrilat), and nitrates.

5HT_{2C} modulators could be useful in treating other diseases associated with obesity, including sleep disorders. Therefore, the compounds described in the present invention could be used in combination with therapeutics for treating sleep disorders.

Examples of suitable therapies for treatment of sleeping disorders for use in combination with the compounds of the present invention include melatonin analogs, melatonin receptor agonists, ML 1 B agonists. GABA A receptor agonists such as barbiturates (e.g., amobarbital, aprobarbital, butabarbital, mephobarbital, pentobarbital, phenobarbital, secobarbital and talbutal), benzodiazepines (e.g., diazepam, lorazepam, oxazepam, alprazolam, chlordiazepoxide, clonazepam, chlorazepate, halazepam and prazepam), also specifically including triazolam

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(Halcion). Other agents for treating sleep disorders include zolpidem (Ambien) and Neurocrine's indiplon..

5HT_{2C} modulators may reduce or ameliorate substance abuse or addictive disorders. Therefore, combination of 5HT_{2C} modulators with agents used to treat addictive disorders may reduce the dose requirement or improve the efficacy of current addictive disorder therapeutics. Examples of agents used to treat substance abuse or addictive disorders are: selective serotonin reuptake inhibitors (SSRI), methadone, buprenorphine, nicotine and bupropion and opiate antagonists.

5HT_{2C} modulators may reduce anxiety or depression; therefore, the compounds described in this application may be used in combination with antianxiety agents or antidepressants. Examples of suitable anti-anxiety agents for use in combination with the compounds of the present invention include benzodiazepines (e.g., diazepam, lorazepam, oxazepam, alprazolam, chlordiazepoxide, clonazepam, chlorazepate, halazepam and prazepam), 5HT_{1A} receptor agonists (e.g., buspirone, flesinoxan, gepirone, ipsapirone and serzone), corticotropin releasing factor (CRF) antagonists and SSRI's.

Examples of suitable classes of anti-depressants for use in combination with the compounds of the present invention include norepinephrine reuptake inhibitors (tertiary and secondary amine tricyclics), selective serotonin reuptake inhibitors (SSRIs) (fluoxetine, fluvoxamine, paroxetine, citalopram and sertraline), monoamine oxidase inhibitors (MAOIs) (isocarboxazid, phenelzine, tranylcypromine, selegiline), reversible inhibitors of monoamine oxidase (RIMAs) (moclobemide), serotonin and norepinephrine reuptake inhibitors (SNRIs) (venlafaxine), corticotropin releasing factor (CRF) receptor antagonists (Britsol-Myers Squibb U.S. patents 6,642,230; 6,630,476; 6,589,952; 6,579,876; 6,525,056; 6,521,636; 6,518,271; 6,515,005; 6,448,261; 6,399,609; 6,362,180; and 6,358,950), alpha-adrenoreceptor antagonists, and atypical antidepressants (bupropion, lithium, nefazodone, trazodone and viloxazine).

The combination of a conventional antipsychotic drug with a $5HT_{2C}$ modulator could also enhance symptom reduction in the treatment of psychosis or mania. Further, such a combination could enable rapid symptom reduction, reducing the need for chronic treatment with antipsychotic agents. Such a combination could also

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reduce the effective antipsychotic dose requirement, resulting in reduced probability of developing the motor dysfunction typical of chronic antipsychotic treatment.

Examples of suitable antipsychotic agents for use in combination with the compounds of the present invention include the phenothiazine (chlorpromazine, mesoridazine, thioridazine, acetophenazine, fluphenazine, perphenazine and trifluoperazine), thioxanthine (chlorprothixene, thiothixene), heterocyclic dibenzazepine (clozapine, olanzepine and aripiprazole), butyrophenone (haloperidol), diphenylbutylpiperidine (pimozide) and indolone (molindolone) classes of antipsychotic agents. Other antipsychotic agents with potential therapeutic value in combination with the compounds in the present invention include loxapine, sulpiride and risperidone.

Combination of the compounds in the present invention with conventional antipsychotic drugs could also provide an enhanced therapeutic effect for the treatment of schizophrenic disorders, as described above for manic disorders. As used here, schizophrenic disorders include paranoid, disorganized, catatonic, undifferentiated and residual schizophrenia, schizophreniform disorder, schizoaffective disorder, delusional disorder, brief psychotic disorder and psychotic disorder not specified. Examples of suitable antipsychotic drugs for combination with the compounds in the present invention include the antipsychotics mentioned above, as well as dopamine receptor antagonists, muscarinic receptor agonists, 5HT_{2A} receptor antagonists and 5HT_{2A}/dopamine receptor antagonists or partial agonists (e.g., olanzepine, aripiprazole, risperidone, ziprasidone).

The compounds described in the present invention could be used to enhance the effects of cognition-enhancing agents, such as acetylcholinesterase inhibitors (e.g., tacrine the active agent in Cognex®), ADHD agents (e.g. methyl-phenidate, atomoxetine the active agent in Strattera® and histamine 3 antagonists), muscarinic receptor-1 agonists (e.g., milameline), nicotinic agonists, glutamic acid receptor (AMPA and NMDA) modulators such as memantine, and nootropic agents (e.g., piracetam, levetiracetam). Examples of suitable therapies for treatment of Alzheimer's disease and cognitive disorders for use in combination with the compounds of the present invention include donepezil, tacrine, revastigraine, 5HT6 receptor antagonists,

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gamma secretase inhibitors, beta secretase inhibitors, SK channel blockers, Maxi-K blockers, and KCNQs blockers.

The compounds described in the present invention could be used to enhance the effects of agents used in the treatment of Parkinson's Disease. Examples of agents used to treat Parkinson's Disease include: levadopa with or without a COMT inhibitor, antiglutamatergic drugs (amantadine, riluzole), alpha-2 adrenergic antagonists such as idazoxan, opiate antagonists, such as naltrexone, other dopamine agonists or transportor modulators, such as ropinirole, or pramipexole or neurotrophic factors such as glial derived neurotrophic factor (GDNF).

The compounds described in the present invention could be used in combination with agents used to treat erectile dysfunction. Examples of suitable treatment for erectile dysfunction include sildenafil (Viagra), vardenafil (Levitra) and tadalafil (Cialis). Other compounds that could be used in combination for erectile dysfunction include yohimbine, phentolamine and papaverine.

The compounds described in the present invention could be used in combination with suitable anti-inflammatory agents. Examples of suitable antiinflammatory agents for use in combination with the compounds of the present invention include prednisone, dexamethasone, cyclooxygenase inhibitors (i.e., COX-1 and/or COX-2 inhibitors such as NSAIDs, aspirin, indomethacin, ibuprofen, piroxicam, Naproxen®, Celebrex®, Vioxx®, Arcoxia®, and Bextra®), CTLA4-Ig agonists/antagonists, CD40 ligand antagonists, IMPDH inhibitors, such as mycophenolate (CellCept®), integrin antagonists, alpha-4 beta-7 integrin antagonists, cell adhesion inhibitors, interferon gamma antagonists, ICAM-1 inhibitor, tumor necrosis factor (TNF) antagonists (e.g., infliximab, OR1384, including TNF-alpha inhibitors, such as tenidap, anti-TNF antibodies or soluble TNF receptor such as etanercept (Enbrel®), Remicade®, rapamycin (sirolimus or Rapamune) and leflunomide (Arava)), prostaglandin synthesis inhibitors, budesonide, clofazimine, CNI-1493, CD4 antagonists (e.g., priliximab), p38 mitogen-activated protein kinase inhibitors, protein tyrosine kinase (PTK) inhibitors, IKK inhibitors, and therapies for the treatment of irritable bowel syndrome (e.g., Zelnorm® and Maxi-K® openers such as those disclosed in U.S. Patent No. 6,184,231 B1).

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Exemplary of such other therapeutic agents which may be used in combination with 5HT_{2C} modulators include the following: cyclosporins (e.g., cyclosporin A), anti-IL-2 receptor (Anti-Tac), anti-CD45RB, anti-CD2, anti-CD3 (OKT-3), anti-CD4, anti-CD80, anti-CD86, monoclonal antibody OKT3, agents blocking the interaction between CD40 and gp39, such as antibodies specific for CD40 and/or gp39 (i.e., CD154), fusion proteins constructed from CD40 and gp39 (CD40Ig and CD8gp39), inhibitors, such as nuclear translocation inhibitors, of NF-kappa B function, such as deoxyspergualin (DSG), gold compounds, antiproliferative agents such as methotrexate, FK506 (tacrolimus, Prograf), mycophenolate mofetil, cytotoxic drugs such as azathiprine and cyclophosphamide, anticytokines such as antiIL-4 or IL-4 receptor fusion proteins and PDE 4 inhibitors such as Ariflo, and the PTK inhibitors disclosed in the following U.S. patent applications, incorporated herein by reference in their entirety: Ser. No. 09/097,338, filed Jun. 15, 1998; Ser. No. 09/094,797, filed Jun. 15, 1998; Ser. No. 09/173,413, filed Oct. 15, 1998; and Ser. No. 09/262,525, filed Mar. 4, 1999. See also the following documents and references cited therein and incorporated herein by reference: Hollenbaugh, D., Et Al, "Cleavable CD40Ig Fusion Proteins and the Binding to Sgp39", J. Immunol. Methods (Netherlands), 188(1), pp. 1-7 (Dec. 15, 1995); Hollenbaugh, D., et al, "The Human T Cell Antigen Gp39, A Member of the TNF Gene Family, Is a Ligand for the CD40 Receptor: Expression of a Soluble Form of Gp39 with B Cell Co-Stimulatory Activity", EMBO J (England), 11(12), pp. 4313-4321 (December 1992); and Moreland, L. W. et al., "Treatment of Rheumatoid Arthritis with a Recombinant Human Tumor Necrosis Factor Receptor (P75)-Fc Fusion Protein," New England J. of Medicine, 337(3), pp. 141-147 (1997).

The above other therapeutic agents, when employed in combination with the compounds of the present invention, may be used, for example, in those amounts indicated in the Physicians' Desk Reference (PDR) or as otherwise determined by one of ordinary skill in the art.

The compounds of formula I of the invention can be administered orally or parenterally, such as subcutaneously or intravenously, as well as by nasal application, transdermally, rectally or sublingually to various mammalian species known to be subject to such maladies, e.g., humans, in an effective amount within the dosage range

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of about 0.2 to 1000 mg, preferably from about 1 to 100 mg in a regimen of single, two or four divided daily doses.

The compounds of the formula I can be administered for any of the uses described herein by any suitable means, for example, orally, such as in the form of tablets, capsules, granules or powders; sublingually; bucally; parenterally, such as by subcutaneous, intravenous, intramuscular, or intracisternal injection or infusion techniques (e.g., as sterile injectable aqueous or non-aqueous solutions or suspensions); nasally, including administration to the nasal membranes, such as by inhalation spray; topically, such as in the form of a cream or ointment; or rectally such as in the form of suppositories; in dosage unit formulations containing non-toxic, pharmaceutically acceptable vehicles or diluents. The present compounds can, for example, be administered in a form suitable for immediate release or extended release. Immediate release or extended release can be achieved by the use of suitable pharmaceutical compositions comprising the present compounds, or, particularly in the case of extended release, by the use of devices such as subcutaneous implants or osmotic pumps. The present compounds can also be administered liposomally.

Exemplary compositions for oral administration include suspensions which can contain, for example, microcrystalline cellulose for imparting bulk, alginic acid or sodium alginate as a suspending agent, methylcellulose as a viscosity enhancer, and sweeteners or flavoring agents such as those known in the art; and immediate release tablets which can contain, for example, microcrystalline cellulose, dicalcium phosphate, starch, magnesium stearate and/or lactose and/or other excipients, binders, extenders, disintegrants, diluents and lubricants such as those known in the art. The compounds of formula I can also be delivered through the oral cavity by sublingual and/or buccal administration. Molded tablets, compressed tablets or freeze-dried tablets are exemplary forms which may be used. Exemplary compositions include those formulating the present compound(s) with fast dissolving diluents such as mannitol, lactose, sucrose and/or cyclodextrins. Also included in such formulations may be high molecular weight excipients such as celluloses (avicel) or polyethylene glycols (PEG). Such formulations can also include an excipient to aid mucosal adhesion such as hydroxy propyl cellulose (HPC), hydroxy propyl methyl cellulose (HPMC), sodium carboxy methyl cellulose (SCMC), maleic anhydride copolymer

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(e.g., Gantrez), and agents to control release such as polyacrylic copolymer (e.g. Carbopol 934). Lubricants, glidants, flavors, coloring agents and stabilizers may also be added for ease of fabrication and use.

Exemplary compositions for nasal aerosol or inhalation administration include solutions in saline which can contain, for example, benzyl alcohol or other suitable preservatives, absorption promoters to enhance bioavailability, and/or other solubilizing or dispersing agents such as those known in the art.

Exemplary compositions for parenteral administration include injectable solutions or suspensions which can contain, for example, suitable non-toxic, parenterally acceptable diluents or solvents, such as mannitol, 1,3-butanediol, water, Ringer's solution, an isotonic sodium chloride solution, or other suitable dispersing or wetting and suspending agents, including synthetic mono- or diglycerides, and fatty acids, including oleic acid, or Cremaphor.

Exemplary compositions for rectal administration include suppositories which can contain, for example, a suitable non-irritating excipient, such as cocoa butter, synthetic glyceride esters or polyethylene glycols, which are solid at ordinary temperatures, but liquify and/or dissolve in the rectal cavity to release the drug.

Exemplary compositions for topical administration include a topical carrier such as Plastibase (mineral oil gelled with polyethylene).

It will be understood that the specific dose level and frequency of dosage for any particular subject can be varied and will depend upon a variety of factors including the activity of the specific compound employed, the metabolic stability and length of action of that compound, the species, age, body weight, general health, sex and diet of the subject, the mode and time of administration, rate of excretion, drug combination, and severity of the particular condition.

PHARMACOLOGICAL ANALYSIS

The pharmacological analysis of each compound for either antogonism or agonism of at 5-HT_{2A}, 5-HT_{2B} and 5-HT_{2C} receptors consisted of in vitro and in vivo studies. In vitro analyses included K_i determinations at 5-HT_{2A}, 5-HT_{2B} and 5-HT_{2C} receptors and an assessment of functional (i.e., agonism or antagonism) activity at each receptor class by IP3 hydrolysis assays. Additional receptor assays were

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conducted to evaluate receptor specificity of 5-HT_{2C} receptors over monoamine and nuisance receptors (e.g. histamine, dopamine, and muscarinic). A compound is considered active as a 5-HT_{2C} agonist if it has an IC₅₀ value or a K_i value of less than about 50 micromolar; preferably less than about 1.0 micromolar; more preferably less than about 0.1 micromolar. Using the assays disclosed herein, compounds of the present invention have been shown to have an IC₅₀ value of less than about 50 micromolar for 5-HT_{2C} agonism.

In vivo assays assessed compound activity in a variety of behavioral paradigms including acute and chronic feeding models, anxiety and depression models (learned-helplessness, elevated plus maze, Geller-Siefter, conditioned taste aversion, taste reactivity, satiety sequence). In aggregate, these models reflect activity as a 5-HT_{2C} agonist (feeding models, anxiety models, depression models) and provide some indication as to bioavailability, metabolism and pharmacokinetics.

Radioligand binding experiments were conducted on recombinant human 5-HT_{2A}, 5-HT_{2B}, and 5-HT_{2C} receptors expressed in HEK293E cells. The affinities of compounds of the present invention to bind at these receptors is determined by their capacity to compete for [¹²⁵I]-1-(2,5-dimethoxy-4-iodophenyl)-2-amino-propane (DOI) binding at the 5-HT_{2A}, 5-HT_{2B}, or 5-HT_{2C}. General references for binding assays include 1) Lucaites VL, Nelson DL, Wainscott DB, Baez M (1996) Receptor subtype and density determine the coupling repertoire of the 5-HT₂ receptor subfamily. Life Sci., 59(13):1081-95. J Med Chem 1988 Jan;31(1):5-7; 2) Glennon RA, Seggel MR, Soine WH, Herrick-Davis K, Lyon RA, Titeler M (1988) [125I]-1-(2,5-dimethoxy-4-iodophenyl)-2-amino-propane: an iodinated radioligand that specifically labels the agonist high-affinity state of 5-HT2 serotonin receptors. J Med. Chem. 31(1):5-7 and 3) Leonhardt S, Gorospe E, Hoffman BJ, Teitler M (1992) Molecular pharmacological differences in the interaction of serotonin with 5-hydroxytryptamine1C and 5-hydroxytryptamine2 receptors. Mol Pharmacol., 42(2):328-35.

The functional properties of compounds (efficacy and potency) were determined in whole cells expressing 5-HT_{2A}, 5-HT_{2B}, or 5-HT_{2C} receptors by assessing their ability to stimulate or inhibit receptor-mediated phosphoinositol hydrolysis. The procedures used are described below.

IN VITRO BINDING ASSAYS

Stable Expression of 5-HT_{2A}, 5-HT_{2B} and 5-HT_{2C} Receptors in HEK293E Cells

Stable cell lines were generated by transfecting 293EBNA cells with plasmids

containing human 5-HT_{2A}, 5-HT_{2B}, or 5-HT_{2C} receptor (INI, INV, VNV or VGV

RNA-edited isoforms) cDNA using calcium phosphate. These plasmids also

contained the cytomegalovirus (CMV) immediate early promoter to drive receptor

expression and EBV oriP for their maintenance as an extrachromosomal element, and
the hph gene from E. Coli to yield hygromycin B resistance (Horlick et al., 1997).

Transfected cells were maintained in Dulbecco's Modified Eagle medium (DMEM)

containing dialyzed 10% fetal bovine serum at 37°C in a humid environment (5%

CO₂) for 10 days. The 5-HT_{2A} cells were adapted to spinner culture for bulk
processing whereas it was necessary to maintain the other lines as adherent cultures.

On the day of harvest, cells were washed in phosphate-buffered saline (PBS),

Membrane Preparation

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On the day of assay, pellets of whole cells (containing approximately 1 X 10⁸ cells) expressing the 5-HT_{2A}, 5-HT_{2B} or 5-HT_{2C} receptor were thawed on ice and homogenized in 50 mM Tris HCl (pH 7.7) containing 1.0 mM EDTA using a Brinkman Polytron (PT-10, setting 6 for 10 sec). The homogenate was centrifuged at 48,000 x g for 10 min and the resulting pellet washed twice by repeated homogenization and centrifugation steps. The final pellet was resuspended in tissue buffer and protein determinations were made by the bichichoninic acid (BCA) assay (Pierce Co., IL) using bovine serum albumin as the standard.

Radioligand Binding Assays for the 5-HT_{2A}, 5-HT_{2B} and 5-HT_{2C} Receptors

Radioligand binding studies were conducted to determine the binding affinities (Ki values) of compounds for the human recombinant 5-HT_{2A}, 5-HT_{2B}, and 5-HT_{2C} receptors (Fitzgerald et al., 1999). Assays were conducted in disposable polypropylene 96-well plates (Costar Corp., Cambridge, MA) and were initiated by the addition of 5-HT_{2A}, 5-HT_{2B}, or 5-HT_{2C} membrane homogenate in tissue buffer

(10-30 (g/well) to assay buffer (50 mM Tris HCl, 0.5 mM EDTA, 10 mM pargyline, 10 mM MgSO₄, 0.05% ascorbic acid, pH 7.5) containing [¹²⁵I]DOI for the 5-HT_{2A} and 5-HT_{2C} receptors (0.3-0.5 nM, final) or [³H]LSD (1-2.0 nM, final) for the 5-HT_{2B} receptor, with or without competing drug (i.e, newly synthesized chemical entity). For a typical competition experiment, a fixed concentration of radioligand was competed with duplicate concentrations of ligand (12 concentrations ranging from 10 picomolar to 10 micromolar). The reaction mixtures were incubated to equilibrium for 45 min at 37°C and terminated by rapid filtration (Packard cell harvester; Perkin-

polyethyleneimine. Filters were washed in ice-cold 50 mM Tris HCl buffer (pH 7.5) and then counted on a Top Count (Packard).

Elmer) over GFF glass-fiber filters that had been pre-soaked in 0.3%

Phosphoinositide Hydrolysis Studies

The ability of newly synthesized compounds to stimulate phosphoinositide 15 (PI) hydrolysis was monitored in whole cells using a variant (Egan et al., 1998) of a protocol described previously (Berridge et al., 1982). HEK293E cells expressing the human 5-HT_{2A}, 5-HT_{2B}, or 5-HT_{2C} receptor were lifted with 0.5 mM EDTA and plated at a density of 100,000/well onto poly-D-lysine-coated 24-well plates (Biocoat; Becton Dickinson, Bedford, MA) in Dulbecco's modified Eagle's serum (DMEM; 20 Gibco BRL) containing high glucose, 2mM glutamine, 10% dialyzed fetal calf serum, 250 (g/ml hygromycin B, and 250(g/ml G418. Following a 24-48 hr period, the growth media was removed and replaced with DMEM without fetal calf serum and inositol (Gibco BRL). The cells were then incubated with DMEM (without serum and inositol) containing a final concentration of 0.5 uCi/well myo-[3H]inositol for 16-25 18 hr. Following this incubation, the cells were washed with DMEM (without serum or inositol) containing 10 mM LiCl and 10 (M pargyline and then incubated for 30 min with the same media but now containing one of several test compounds. Reactions were terminated by aspirating the media and lysing the cells by freezethaw. [3H]phosphoinositides were extracted with chloroform/methanol (1:2 v/v), 30 separated by anion exchange chromatography (Bio-Rad AGI-X8 resin), and counted by liquid scintillation spectroscopy as described previously (Egan et al., 1998).

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Calcium Fluorescence Studies

The ability of newly synthesized compounds to stimulate calcium fluorescence was monitored in whole cells using a protocol described previously (Fitzgerlad et al., 1999). HEK293E cells expressing the human 5-HT_{2C}, or 5-HT_{2B} receptor were lifted with 0.5 mM EDTA and plated at a density of 50,000/well onto poly-D-lysine-coated 96-well plates (Biocoat; Becton Dickinson, Bedford, MA) in Dulbecco's modified Eagle's serum (DMEM; Gibco BRL) containing high glucose, 2mM glutamine, 10% dialyzed fetal calf serum, 250 μg/ml hygromycin B, and 250 μg/ml G418. Following a 24 hr period, the cell plates are removed from the incubator and an equal volume of Loading Buffer (Hanks BSS with 200mM HEPES, pH 5.98) containing the calcium dye reagent (Fluo-3) is added to each well (100 μL per well for 96-well plates and then incubated for 1 hour at 37C. Following the dye loading of the cells he plates are transferred to the FLIPR. Test compounds are added to the plate as a concentration response curve and the changes in fluorescence units due to calcium influx are monitored for a period of three seconds.

Data Analyses

The equilibrium apparent dissociation constants (Ki's) from the competition experiments were calculated using an iterative nonlinear regression curve-fitting program (Excelfit and TA Activity Base). For the PI hydrolysis and FLIPR experiments, EC50's were calculated using a one-site 'pseudo' Hill model: y=((Rmax-Rmin)/(1+R/EC50)nH)) + Rmax where R= response (GraphPad Prism; San Diego, CA). Emax (maximal response) was derived from the fitted curve maxima (net IP stimulation) for each compound. Intrinsic activity (IA) was determined by expressing the Emax of a compound as a percentage of the Emax of 5-HT (IA=1.0).

Efficacy Models to Evaluate Food Consumption and Weight Loss

Acute overnight feeding assay. Compounds are assessed to for their ability to reduce food consumption during the dark cycle, which is the most active period of feeding in the rat. Fischer 344 rats are trained on a fixed ratio three (FR3) response paradigm which requires them to press a bar 3 consecutive times in order to obtain a food pellet. The number of bar presses occurring throughout the dark cycle can be

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monitored electronically as a measure of food intake by the animal. Rats are dosed orally or intraperitoneally with test compound 30 minutes prior to the onset of the dark cycle. The treated animals are then placed in individual operant boxes for 15 hours (12 hrs of dark cycle and the first three hours of the light cycle). Food intake in compound treated animals is compared to that of vehicle treated animals in order to determine percent reductions in food intake. Simultaneous measurements of water intake and locomotor activity are also measured during the period to assess for potential adverse effects.

10 Chronic Feeding Assay

Compounds are assessed for their long term impact on food intake and body weight in a three to fourteen week chronic treatment paradigm in Sprague-Dawley rats (starting weight ~450 g). Male Sprague-Dawley rats are pre-handled for one week prior to the onset of dosing during which time they are also assessed for food intake behavior. Rats are then assigned to treatment groups. Rats are dosed with vehicle or compound by oral gavage. The food intake and body weights are cumulatively assessed at the end of each treatment week and compared to vehicle treated animals. In some studies food intake is measured daily in order to assess the impact of reduced food consumption on pair-fed animals. At the end of the study period the animals are assessed for changes in body composition utilizing DEXA and are then sacrificed in order to examine changes in various blood plasma parameters.

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DOSAGE AND FORMULATIONS

The serotonin agonist and serotonin antagonist compounds of this invention can be administered as treatment for the control or prevention of central nervous system disorders including obesity, anxiety, depression, psychosis, schizophrenia, sleep and sexual disorders, migraine and other conditions associated with cephalic pain, social phobias, and gastrointestinal disorders such as dysfunction of the gastrointestinal tract motility by any means that produces contact of the active agent with the agent's site of action, i.e., 5-HT2 receptors, in the body of a mammal. It can be administered by any conventional means available for use in conjunction with pharmaceuticals, either as an individual therapeutic agent or in a combination of therapeutic agents. It can be administered alone, but preferably is administered with a pharmaceutical carrier selected on the basis of the chosen route of administration and standard pharmaceutical practice.

The compounds of the present invention can be administered in such oral dosage forms as tablets, capsules (each of which includes sustained release or timed release formulations), pills, powders, granules, elixirs, tinctures, suspensions, syrups, and emulsions. Likewise, they may also be administered in intravenous (bolus or infusion), intraperitoneal, subcutaneous, or intramuscular form, all using dosage forms well known to those of ordinary skill in the pharmaceutical arts.

The dosage administered will, of course, vary depending upon known factors, such as the pharmacodynamic characteristics of the particular agent and its mode and route of administration; the age, health and weight of the recipient; the nature and

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extent of the symptoms; the kind of concurrent treatment; the frequency of treatment; and the effect desired. By way of general guidance, a daily dosage of active ingredient can be expected to be about 0.001 to about 1000 milligrams per kilogram of body weight, with the preferred dose being about 0.01 to about 100 mg/kg; with the more preferred dose being about 0.01 to about 30 mg/kg. Advantageously, compounds of the present invention may be administered in a single daily dose, or the total daily dosage may be administered in divided doses of two, three, or four times daily.

Dosage forms of compositions suitable for administration contain from about 0.5 mg to about 100 mg of active ingredient per unit. In these pharmaceutical compositions the active ingredient will ordinarily be present in an amount of about 0.5-95% by weight based on the total weight of the composition. The active ingredient can be administered orally in solid dosage forms, such as capsules, tablets and powders, or in liquid dosage forms, such as elixirs, syrups and suspensions. It can also be administered parenterally, in sterile liquid dosage forms.

Gelatin capsules contain the active ingredient and powdered carriers, such as lactose, starch, cellulose derivatives, magnesium stearate, stearic acid, and the like. Similar diluents can be used to make compressed tablets. Both tablets and capsules can be manufactured as sustained release products to provide for continuous release of medication over a period of hours. Compressed tablets can be sugar coated or film coated to mask any unpleasant taste and protect the tablet from the atmosphere, or enteric coated for selective disintegration in the gastrointestinal tract. Liquid dosage forms for oral administration can contain coloring and flavoring to increase patient acceptance.

In general, water, a suitable oil, saline, aqueous dextrose (glucose), and related sugar solutions and glycols such as propylene glycol or polyethylene glycols are suitable carriers for parenteral solutions. Solutions for parenteral administration preferably contain a water soluble salt of the active ingredient, suitable stabilizing agents, and if necessary, buffer substances. Antioxidizing agents such as sodium bisulfite, sodium sulfite, or ascorbic acid, either alone or combined, are suitable stabilizing agents. Also used are citric acid and its salts, and sodium EDTA. In addition, parenteral solutions can contain preservatives, such as benzalkonium

chloride, methyl- or propyl-paraben and chlorobutanol. Suitable pharmaceutical carriers are described in *Remington's Pharmaceutical Sciences*, *supra*, a standard reference text in this field.

Useful pharmaceutical dosage-forms for administration of the compounds of this invention can be illustrated as follows:

Capsules

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A large number of unit capsules can be prepared by filling standard two-piece hard gelatin capsules each with 100 mg of powdered active ingredient, 150 mg of lactose, 50 mg of cellulose, and 6 mg magnesium stearic.

Soft Gelatin Capsules

A mixture of active ingredient in a digestible oil such as soybean oil, cottonseed oil or olive oil can be prepared and injected by means of a positive displacement pump into gelatin to form soft gelatin capsules containing 100 mg of the active ingredient. The capsules should then be washed and dried.

Tablets

A large number of tablets can be prepared by conventional procedures so that the dosage unit is 100 mg of active ingredient, 0.2 mg of colloidal silicon dioxide, 5 milligrams of magnesium stearate, 275 mg of microcrystalline cellulose, 11 mg of starch and 98.8 mg of lactose. Appropriate coatings may be applied to increase palatability or delay absorption.

25 Suspension

An aqueous suspension can be prepared for oral administration so that each 5 mL contain 25 mg of finely divided active ingredient, 200 mg of sodium carboxymethyl cellulose, 5 mg of sodium benzoate, 1.0 g of sorbitol solution, U.S.P., and 0.025 mg of vanillin.

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Injectable

A parenteral composition suitable for administration by injection can be prepared by stirring 1.5% by weight of active ingredient in 10% by volume propylene glycol and water. The solution is sterilized by commonly used techniques.

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While it is apparent that the embodiments of the invention herein disclosed are well suited to fulfill the objectives stated above, it will be appreciated that numerous modifications and other embodiments may be implemented by those skilled in the art, and it is intended that the appended claims cover all such modifications and embodiments that fall within the true spirit and scope of the present invention.

A number of references have been cited and the entire disclosures of which are incorporated herein by reference.